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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6: WO 99/50454 (11) International Publication Number: C12Q 1/68 **A2** (43) International Publication Date: 7 October 1999 (07.10.99) (74) Agents: GRANAHAN, Patricia et al.; Hamilton, Brook, Smith (21) International Application Number: PCT/US99/06473 & Reynolds, P.C., Two Militia Drive, Lexington, MA 02421 (22) International Filing Date: 26 March 1999 (26.03.99) (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, (30) Priority Data: 09/054,272 1 April 1998 (01.04.98) US BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, (63) Related by Continuation (CON) or Continuation-in-Part SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, (CIP) to Earlier Application ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, 09/054,272 (CIP) US 1 April 1998 (01.04.98) UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, Filed on RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, (71) Applicant (for all designated States except US): WHITEHEAD INSTITUTE FOR BIOMEDICAL RESEARCH [US/US]; NE, SN, TD, TG). Nine Cambridge Center, Cambridge, MA 02142 (US). **Published** (72) Inventors; and (75) Inventors/Applicants (for US only): LANDER, Eric, S. Without international search report and to be republished [US/US]; 151 Bishop Allen Drive, Cambridge, MA 02138 upon receipt of that report. (US). DALEY, George, Q. [US/US]; 50 Young Road, Weston, MA 02193 (US). CARGILL, Michele [US/US]; 50 Follen Street #208, Cambridge, MA 02138 (US). IRELAND, James, S. [US/US]; 36 College Avenue #1, Somerville, MA 02144 (US). ROZEN, Steven, G. [US/US]; 45 Josephine Avenue, Somerville, MA 02144-2312 (US).

(54) Title: CODING SEQUENCE POLYMORPHISMS IN VASCULAR PATHOLOGY GENES

(57) Abstract

The invention provides nucleic acid segments of the human genome, particularly nucleic acid segments from the coding region of a gene, including polymorphic sites. Allele-specific primers and probes hybridizing to regions flanking or containing these sites are also provided. The nucleic acids, primers and probes are used in applications such as phenotype correlations, forensics, paternity testing, medicine and genetic analysis.

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CODING SEQUENCE POLYMORPHISMS IN VASCULAR PATHOLOGY GENES

RELATED APPLICATIONS

This application is a Continuation-in-Part of U.S. Application No. 09/054,272, 5 filed April 1, 1998, the contents of which are incorporated herein in their entirety by reference.

BACKGROUND OF THE INVENTION

The genomes of all organisms undergo spontaneous mutation in the course of their continuing evolution, generating variant forms of progenitor sequences (Gusella, 10 Ann. Rev. Biochem. 55, 831-854 (1986)). The variant form may confer an evolutionary advantage or disadvantage relative to a progenitor form or may be neutral. In some instances, a variant form confers a lethal disadvantage and is not transmitted to subsequent generations of the organism. In other instances, a variant form confers an evolutionary advantage to the species and is eventually incorporated 15 into the DNA of many or most members of the species and effectively becomes the progenitor form. In many instances, both progenitor and variant form(s) survive and co-exist in a species population. The coexistence of multiple forms of a sequence gives rise to polymorphisms.

Several different types of polymorphism have been reported. A restriction 20 fragment length polymorphism (RFLP) Is a variation in DNA sequence that alters the length of a restriction fragment (Botstein et al., Am. J. Hum. Genet. 32, 314-331 (1980)). The restriction fragment length polymorphism may create or delete a restriction site, thus changing the length of the restriction fragment. RFLPs have been widely used in human and animal genetic analyses (see WO 90/13668; W090/11369;

25 Donis-Keller, Cell 51, 319-337 (1987); Lander et al., Genetics 121, 85-99 (1989)). When a heritable trait can be linked to a particular RFLP, the presence of the RFLP in an individual can be used to predict the likelihood that the animal will also exhibit the trait.

Other polymorphisms take the form of short tandem repeats (STRs) that 30 include tandem di-, tri- and tetra-nucleotide repeated motifs. These tandem repeats are also referred to as variable number tandem repeat (VNTR) polymorphisms. VNTRs have been used in identity and paternity analysis (US 5,075,217; Armour et al., FEBS Lett. 307, 113-115 (1992); Horn et al., W0 91/14003; Jeffreys, EP 370,719), and in a large number of genetic mapping studies.

Other polymorphisms take the form of single nucleotide variations between individuals of the same species. Such polymorphisms are far more frequent than RFLPs, STRs and VNTRs. Some single nucleotide polymorphisms (SNP) occur in protein-coding sequences (coding sequence SNP (cSNP)), in which case, one of the polymorphic forms may give rise to the expression of a defective or otherwise variant protein and, potentially, a genetic disease. Examples of genes in which polymorphisms within coding sequences give rise to genetic disease include β-globin (sickle cell anemia), apoE4 (Alzheimer's Disease), Factor V Leiden (thrombosis), and CFTR (cystic fibrosis). cSNPs can alter the codon sequence of the gene and therefore specify an alternative amino acid. Such changes are called "missense" when another amino acid is substituted, and "nonsense" when the alternative codon specifies a stop signal in protein translation. When the cSNP does not alter the amino acid specified the cSNP is called "silent".

Other single nucleotide polymorphisms occur in noncoding regions. Some of these polymorphisms may also result in defective protein expression (e.g., as a result 20 of defective splicing). Other single nucleotide polymorphisms have no phenotypic effects.

Single nucleotide polymorphisms can be used in the same manner as RFLPs and VNTRs, but offer several advantages. Single nucleotide polymorphisms occur with greater frequency and are spaced more uniformly throughout the genome than other forms of polymorphism. The greater frequency and uniformity of single nucleotide polymorphisms means that there is a greater probability that such a polymorphism will be found in close proximity to a genetic locus of interest than would be the case for other polymorphisms. The different forms of characterized single nucleotide polymorphisms are often easier to distinguish than other types of polymorphism (e.g., by use of assays employing allele-specific hybridization probes or primers).

Only a small percentage of the total repository of polymorphisms in humans and other organisms has been identified. The limited number of polymorphisms identified to date is due to the large amount of work required for their detection by conventional methods. For example, a conventional approach to identifying polymorphisms might be to sequence the same stretch of DNA in a population of individuals by dideoxy sequencing. In this type of approach, the amount of work

increases in proportion to both the length of sequence and the number of individuals in a population and becomes impractical for large stretches of DNA or large numbers of persons.

SUMMARY OF THE INVENTION

Work described herein pertains to the identification of polymorphisms which can predispose individuals to disease, particularly vascular pathologies, by resequencing large numbers of genes in a large number of individuals. Eighteen genes in a minimum of 30 individuals have been resequenced as described herein, and 92 SNPs have been discovered (see the Table). Forty of these SNPs are cSNPs which specify a different amino acid sequence, while 49 of the SNPs are silent cSNPs. Three of the SNPs were located in non-coding regions.

The invention relates to a gene which comprises a single nucleotide polymorphism at a specific location. In a particular embodiment the invention relates to the variant allele of a gene having a single nucleotide polymorphism, which variant allele differs from a reference allele by one nucleotide at the site(s) identified in the Table. Complements of these nucleic acid segments are also included. The segments can be DNA or RNA, and can be double- or single-stranded. Segments can be, for example, 5-10, 5-15, 10-20, 5-25, 10-30, 10-50 or 10-100 bases long.

The invention further provides allele-specific oligonucleotides that hybridize to a gene comprising a single nucleotide polymorphism or to the complement of the gene. These oligonucleotides can be probes or primers.

The invention further provides a method of analyzing a nucleic acid from an individual. The method determines which base is present at any one of the polymorphic sites shown in the Table. Optionally, a set of bases occupying a set of the polymorphic sites shown in the Table is determined. This type of analysis can be performed on a number of individuals, who are tested for the presence of a disease phenotype. The presence or absence of disease phenotype is then correlated with a base or set of bases present at the polymorphic site or sites in the individuals tested.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-1C are a table illustrating the locations of single nucleotide polymorphisms of various genes.

Figure 2 is a listing of the genes from Figures 1A-C with their corresponding GenBank Accession numbers and the nucleotide position within that sequence at which the single nucleotide polymorphism is located.

Figures 3A-B are a listing of the nucleotide sequence corresponding to GenBank Accession number D10202 for the gene PTAFR.

Figures 4A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number D29832 for the gene AT3.

Figures 5A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number D38081 for the gene TBXA2R.

Figures 6A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number J02703 for the gene ITGB3.

Figures 7A-C are a listing of the nucleotide sequence corresponding to the 10 GenBank Accession number J02764 for the gene ITGA2B.

Figures 8A-F are a listing of the nucleotide sequence corresponding to the GenBank Accession number J02846 for the gene F3.

Figures 9A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number J02898 for the gene CETP.

Figures 10A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number J03225 for the gene TFPI.

Figures 11A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number K02059 for the gene PROC.

Figure 12 is a listing of the nucleotide sequence corresponding to the GenBank 20 Accession number L00336 for the gene LDLR.

Figure 13 is a listing of the nucleotide sequence corresponding to the GenBank Accession number L00338.

Figure 14 is a listing of the nucleotide sequence corresponding to the GenBank Accession number L00343 for the gene LDLR.

Figure 15 is a listing of the nucleotide sequence corresponding to the GenBank Accession number L00344 for the gene LDLR.

Figure 16 is a listing of the nucleotide sequence corresponding to the GenBank Accession number L00345 for the gene LDLR.

Figure 17 is a listing of the nucleotide sequence corresponding to the GenBank 30 Accession number L00347 for the gene LDLR.

Figure 18 is a listing of the nucleotide sequence corresponding to the GenBank Accession number L00349 for the gene LDLR.

Figures 19A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number L00351 for the gene LDLR.

Figures 20A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number L29401 for the gene LDLR.

Figures 21A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number L32765 for the gene F5.

Figures 22A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number M11058 for the gene HMGCR.

Figures 23A-F are a listing of the nucleotide sequence corresponding to the GenBank Accession number M11228 for the gene PROC.

Figures 24A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number M12625 for the gene LCAT.

Figures 25A-C are a listing of the nucleotide sequence corresponding to the 10 GenBank Accession number M12849 for the gene HCF2.

Figures 26A-E are a listing of the nucleotide sequence corresponding to the GenBank Accession number M14335 for the gene F5.

Figures 27A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number M15856 for the gene LPL.

Figures 28A-N are a listing of the nucleotide sequence corresponding to the GenBank Accession number M17262 for the gene F2.

Figures 29A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number M20311 for the gene ITGB3.

Figure 30 is a listing of the nucleotide sequence corresponding to the GenBank 20 Accession number M21645 for the gene AT3.

Figures 31A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number M22569 for the gene ITGA2B.

Figures 32A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number M30185 for the gene CETP.

Figures 33A-H are a listing of the nucleotide sequence corresponding to the GenBank Accession number M33320 for the gene ITGA2B.

Figures 34A-G are a listing of the nucleotide sequence corresponding to the GenBank Accession number M58600 for the gene HCF2.

Figures 35A-B are a listing of the nucleotide sequence corresponding to the 30 GenBank Accession number M62424 for the gene F2R.

Figures 36A-C are a listing of the nucleotide sequence corresponding to the GenBank Accession number M76722 for the gene LPL.

Figures 37A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number U59436 for the gene LDLR.

Figures 38A-B are a listing of the nucleotide sequence corresponding to the GenBank Accession number Z22555 for the gene CLanalog.

DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to a gene which comprises a single nucleotide polymorphism (SNP) at a specific location. The gene which includes the SNP has at least two alleles, referred to herein as the reference allele and the variant allele. The 5 reference allele (prototypical or wild type allele) has been designated arbitrarily and typically corresponds to the nucleotide sequence of the gene which has been deposited with GenBank under a given Accession number. The variant allele differs from the reference allele by one nucleotide at the site(s) identified in the Table. The present invention also relates to variant alleles of the described genes and to complements of 10 the variant alleles. The invention further relates to portions of the variant alleles and portions of complements of the variant alleles which comprise (encompass) the site of the SNP and are at least 5 nucleotides in length. Portions can be, for example, 5-10, 5-15, 10-20, 5-25, 10-30, 10-50 or 10-100 bases long. For example, a portion of a variant allele which is 5 nucleotides in length includes the single nucleotide 15 polymorphism (the nucleotide which differs from the reference allele at that site) and four additional nucleotides which flank the site in the variant allele. These nucleotides can be on one or both sides of the polymorphism. Polymorphisms which are the subject of this invention are defined in the Table with respect to the reference sequence deposited in GenBank under the Accession number indicated. For example, 20 the invention relates to a portion of a gene (e.g., AT3) having a nucleotide sequence as deposited in GenBank (e.g., M21645) comprising a single nucleotide polymorphism at a specific position (e.g., nucleotide 100). The reference allele for AT3 is shown in column 15 and the variant allele is shown in column 17 of the Table. The nucleotide sequences of the invention can be double- or single-stranded.

The invention further provides allele-specific oligonucleotides that hybridize to a gene comprising a single nucleotide polymorphism or to the complement of the gene. These oligonucleotides can be probes or primers.

The invention further provides a method of analyzing a nucleic acid from an individual. The method determines which base is present at any one of the polymorphic sites shown in the Table. Optionally, a set of bases occupying a set of the polymorphic sites shown in the Table is determined. This type of analysis can be performed on a number of individuals, who are tested for the presence of a disease phenotype. The presence or absence of disease phenotype is then correlated with a base or set of bases present at the polymorphic site or sites in the individuals tested.

DEFINITIONS

An oligonucleotide can be DNA or RNA, and single- or double-stranded.

Oligonucleotides can be naturally occurring or synthetic, but are typically prepared by synthetic means. Preferred oligonucleotides of the invention include segments of DNA, or their complements, which include any one of the polymorphic sites shown in the Table. The segments can be between 5 and 250 bases, and, in specific embodiments, are between 5-10, 5-20, 10-20, 10-50, 20-50 or 10-100 bases. The polymorphic site can occur within any position of the segment. The segments can be from any of the allelic forms of DNA shown in the Table.

As used herein, the terms "nucleotide" and "nucleic acid" are intended to be equivalent. The terms "nucleotide sequence", "nucleic acid sequence", "nucleic acid molecule" and "segment" are intended to be equivalent.

Hybridization probes are oligonucleotides which bind in a base-specific manner to a complementary strand of nucleic acid. Such probes include peptide

15 nucleic acids, as described in Nielsen et al., Science 254, 1497-1500 (1991). Probes can be any length suitable for specific hybridization to the target nucleic acid sequence. The most appropriate length of the probe may vary depending upon the hybridization method in which it is being used; for example, particular lengths may be more appropriate for use in microfabricated arrays, while other lengths may be more suitable for use in classical hybridization methods. Suitable probes and primers can range from about 5 nucleotides to about 30 nucleotides in length. For example, probes and primers can be 5, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 25, 26, 28 or 30 nucleotides in length. The probe or primer preferably contains at least one polymorphic site occupied by any of the possible variant nucleotides. The nucleotide sequence can correspond to the coding sequence of the allele or to the complement of the coding sequence of the allele.

As used herein, the term "primer" refers to a single-stranded oligonucleotide which acts as a point of initiation of template-directed DNA synthesis under appropriate conditions (e.g., in the presence of four different nucleoside triphosphates and an agent for polymerization, such as, DNA or RNA polymerase or reverse transcriptase) in an appropriate buffer and at a suitable temperature. The appropriate length of a primer depends on the intended use of the primer, but typically ranges from 15 to 30 nucleotides. Short primer molecules generally require cooler temperatures to form sufficiently stable hybrid complexes with the template. A primer need not reflect the exact sequence of the template, but must be sufficiently complementary to hybridize with a template. The term primer site refers to the area of the target DNA to which a primer hybridizes. The term primer pair refers to a set

of primers including a 5' (upstream) primer that hybridizes with the 5' end of the DNA sequence to be amplified and a 3' (downstream) primer that hybridizes with the complement of the 3' end of the sequence to be amplified.

As used herein, linkage describes the tendency of genes, alleles, loci or genetic 5 markers to be inherited together as a result of their location on the same chromosome. It can be measured by percent recombination between the two genes, alleles, loci or genetic markers.

As used herein, polymorphism refers to the occurrence of two or more genetically determined alternative sequences or alleles in a population. A

10 polymorphic marker or site is the locus at which divergence occurs. Preferred markers have at least two alleles, each occurring at frequency of greater than 1%, and more preferably greater than 10% or 20% of a selected population. A polymorphic locus may be as small as one base pair. Polymorphic markers include restriction fragment length polymorphisms, variable number of tandem repeats (VNTR's),

15 hypervariable regions, minisatellites, dinucleotide repeats, trinucleotide repeats, tetranucleotide repeats, simple sequence repeats, and insertion elements such as Alu. The first identified allelic form is arbitrarily designated as the reference form and other allelic forms are designated as alternative or variant alleles. The allelic form occurring most frequently in a selected population is sometimes referred to as the wildtype form. Diploid organisms may be homozygous or heterozygous for allelic forms. A diallelic or biallelic polymorphism has two forms. A triallelic polymorphism has three forms.

Work described herein pertains to the resequencing of large numbers of genes in a large number of individuals to identify polymorphisms which can predispose individuals to disease, particularly vascular pathologies. Eighteen genes in a minimum of 30 individuals have been resequenced as described herein, and 92 SNPs have been discovered (see the Table). Forty of these SNPs are cSNPs which specify a different amino acid sequence, while 49 of the SNPs are silent cSNPs. Three of the SNPs were located in non-coding regions.

The 18 genes which were subjected to analysis encode proteins that are involved in biochemical pathways that regulate blood coagulation, lipid metabolism, and platelet and endothelial cell function. Polymorphisms in all 18 genes are candidates for genetic factors that influence the pathophysiology of the blood and blood vessels and thus can be relevant to the genetic risk of cardiovascular diseases.

35 The identified polymorphisms can also be relevant to other disease categories.

By altering amino acid sequence, SNPs may alter the function of the encoded proteins. The discovery of the SNP facilitates biochemical analysis of the variants

and the development of assays to characterize the variants and to screen for pharmaceutical that would interact directly with on or another form of the protein. SNPs (including silent SNPs) may also alter the regulation of the gene at the transcriptional or post-transcriptional level. SNPs (including silent SNPs) also enable the development of specific DNA, RNA, or protein-based diagnostics that detect the presence or absence of the polymorphism in particular conditions.

A single nucleotide polymorphism occurs at a polymorphic site occupied by a single nucleotide, which is the site of variation between allelic sequences. The site is usually preceded by and followed by highly conserved sequences of the allele (e.g., sequences that vary in less than 1/100 or 1/1000 members of the populations).

A single nucleotide polymorphism usually arises due to substitution of one nucleotide for another at the polymorphic site. A transition is the replacement of one purine by another purine or one pyrimidine by another pyrimidine. A transversion is the replacement of a purine by a pyrimidine or vice versa. Single nucleotide

15 polymorphisms can also arise from a deletion of a nucleotide or an insertion of a nucleotide relative to a reference allele. Typically the polymorphic site is occupied by a base other than the reference base. For example, where the reference allele contains the base "T" at the polymorphic site, the altered allele can contain a "C", "G" or "A" at the polymorphic site.

Hybridizations are usually performed under stringent conditions, for example, at a salt concentration of no more than 1 M and a temperature of at least 25°C. For example, conditions of 5X SSPE (750 mM NaCl, 50 mM NaPhosphate, 5 mM EDTA, pH 7.4) and a temperature of 25-30°C, or equivalent conditions, are suitable for allele-specific probe hybridizations. Equivalent conditions can be determined by varying one or more of the parameters given as an example, as known in the art, while maintaining a similar degree of identity or similarity between the target nucleotide sequence and the primer or probe used.

The term "isolated" is used herein to indicate that the material in question exists in a physical milieu distinct from that in which it occurs in nature. For example, an isolated nucleic acid of the invention may be substantially isolated with respect to the complex cellular milieu in which it naturally occurs. In some instances, the isolated material will form part of a composition (for example, a crude extract containing other substances), buffer system or reagent mix. In other circumstance, the material may be purified to essential homogeneity, for example as determined by PAGE or column chromatography such as HPLC. Preferably, an isolated nucleic acid comprises at least about 50, 80 or 90 percent (on a molar basis) of all macromolecular species present.

I. Novel Polymorphisms of the Invention

The novel polymorphisms of the invention are shown in the Table.

II. Analysis of Polymorphisms

A. Preparation of Samples

Polymorphisms are detected in a target nucleic acid from an individual being analyzed. For assay of genomic DNA, virtually any biological sample (other than pure red blood cells) is suitable. For example, convenient tissue samples include whole blood, semen, saliva, tears, urine, fecal material, sweat, buccal, skin and hair. For assay of cDNA or mRNA, the tissue sample must be obtained from an organ in which the target nucleic acid is expressed. For example, if the target nucleic acid is a cytochrome P450, the liver is a suitable source.

Many of the methods described below require amplification of DNA from target samples. This can be accomplished by e.g., PCR. See generally PCR Technology: Principles and Applications for DNA Amplification (ed. H.A. Erlich,

- 15 Freeman Press, NY, NY, 1992); PCR Protocols: A Guide to Methods and Applications (eds. Innis, et al., Academic Press, San Diego, CA, 1990); Mattila et al., Nucleic Acids Res. 19, 4967 (1991); Eckert et al., PCR Methods and Applications 1, 17 (1991); PCR (eds. McPherson et al., IRL Press, Oxford); and U.S. Patent 4,683,202.
- Other suitable amplification methods include the ligase chain reaction (LCR) (see Wu and Wallace, *Genomics* 4, 560 (1989), Landegren et al., Science 241, 1077 (1988), transcription amplification (Kwoh et al., Proc. Natl. Acad. Sci. USA 86, 1173 (1989)), and self-sustained sequence replication (Guatelli et al., Proc. Nat. Acad. Sci. USA, 87, 1874 (1990)) and nucleic acid based sequence amplification (NASBA). The
- 25 latter two amplification methods involve isothermal reactions based on isothermal transcription, which produce both single stranded RNA (ssRNA) and double stranded DNA (dsDNA) as the amplification products in a ratio of about 30 or 100 to 1, respectively.

B. Detection of Polymorphisms in Target DNA

- There are two distinct types of analysis of target DNA for detecting polymorphisms. The first type of analysis, sometimes referred to as de novo characterization, is carried out to identify polymorphic sites not previously characterized (i.e., to identify new polymorphisms). This analysis compares target sequences in different individuals to identify points of variation, i.e., polymorphic
- 35 sites. By analyzing groups of individuals representing the greatest ethnic diversity

among humans and greatest breed and species variety in plants and animals, patterns characteristic of the most common alleles/haplotypes of the locus can be identified, and the frequencies of such alleles/haplotypes in the population can be determined. Additional allelic frequencies can be determined for subpopulations characterized by criteria such as geography, race, or gender. The de novo identification of polymorphisms of the invention is described in the Examples section. The second type of analysis determines which form(s) of a characterized (known) polymorphism are present in individuals under test. There are a variety of suitable procedures, which are discussed in turn.

10 1. Allele-Specific Probes

The design and use of allele-specific probes for analyzing polymorphisms is described by e.g., Saiki et al., Nature 324, 163-166 (1986); Dattagupta, EP 235,726, Saiki, WO 89/11548. Allele-specific probes can be designed that hybridize to a segment of target DNA from one individual but do not hybridize to the corresponding segment from another individual due to the presence of different polymorphic forms in the respective segments from the two individuals. Hybridization conditions should be sufficiently stringent that there is a significant difference in hybridization intensity between alleles, and preferably an essentially binary response, whereby a probe hybridizes to only one of the alleles. Some probes are designed to hybridize to a segment of target DNA such that the polymorphic site aligns with a central position (e.g., in a 15-mer at the 7 position; in a 16-mer, at either the 8 or 9 position) of the probe. This design of probe achieves good discrimination in hybridization between different allelic forms.

Allele-specific probes are often used in pairs, one member of a pair showing a perfect match to a reference form of a target sequence and the other member showing a perfect match to a variant form. Several pairs of probes can then be immobilized on the same support for simultaneous analysis of multiple polymorphisms within the same target sequence.

2. Tiling Arrays

The polymorphisms can also be identified by hybridization to nucleic acid arrays, some examples of which are described in WO 95/11995. One form of such arrays is described in the Examples section in connection with de novo identification of polymorphisms. The same array or a different array can be used for analysis of characterized polymorphisms. WO 95/11995 also describes subarrays that are optimized for detection of a variant form of a precharacterized polymorphism. Such a

subarray contains probes designed to be complementary to a second reference sequence, which is an allelic variant of the first reference sequence. The second group of probes is designed by the same principles as described in the Examples, except that the probes exhibit complementarity to the second reference sequence. The inclusion of a second group (or further groups) can be particularly useful for analyzing short subsequences of the primary reference sequence in which multiple mutations are expected to occur within a short distance commensurate with the length of the probes (e.g., two or more mutations within 9 to 21 bases).

3. Allele-Specific Primers

An allele-specific primer hybridizes to a site on target DNA overlapping a polymorphism and only primes amplification of an allelic form to which the primer exhibits perfect complementarity. See Gibbs, *Nucleic Acid Res.* 17, 2427-2448 (1989). This primer is used in conjunction with a second primer which hybridizes at a distal site. Amplification proceeds from the two primers, resulting in a detectable product which indicates the particular allelic form is present. A control is usually performed with a second pair of primers, one of which shows a single base mismatch at the polymorphic site and the other of which exhibits perfect complementarity to a distal site. The single-base mismatch prevents amplification and no detectable product is formed. The method works best when the mismatch is included in the 3'-20 most position of the oligonucleotide aligned with the polymorphism because this position is most destabilizing to elongation from the primer (see, e.g., WO 93/22456).

4. Direct-Sequencing

The direct analysis of the sequence of polymorphisms of the present invention can be accomplished using either the dideoxy chain termination method or the Maxam 25 Gilbert method (see Sambrook et al., Molecular Cloning, A Laboratory Manual (2nd Ed., CSHP, New York 1989); Zyskind et al., Recombinant DNA Laboratory Manual, (Acad. Press, 1988)).

5. Denaturing Gradient Gel Electrophoresis

Amplification products generated using the polymerase chain reaction can be analyzed by the use of denaturing gradient gel electrophoresis. Different alleles can be identified based on the different sequence-dependent melting properties and electrophoretic migration of DNA in solution. Erlich, ed., PCR Technology, Principles and Applications for DNA Amplification, (W.H. Freeman and Co, New York, 1992), Chapter 7.

6. Single-Strand Conformation Polymorphism Analysis

Alleles of target sequences can be differentiated using single-strand conformation polymorphism analysis, which identifies base differences by alteration in electrophoretic migration of single stranded PCR products, as described in Orita et al., Proc. Nat. Acad. Sci. 86, 2766-2770 (1989). Amplified PCR products can be generated as described above, and heated or otherwise denatured, to form single stranded amplification products. Single-stranded nucleic acids may refold or form secondary structures which are partially dependent on the base sequence. The different electrophoretic mobilities of single-stranded amplification products can be related to base-sequence differences between alleles of target sequences.

III. Methods of Use

After determining polymorphic form(s) present in an individual at one or more polymorphic sites, this information can be used in a number of methods.

A. Forensics

Determination of which polymorphic forms occupy a set of polymorphic sites in an individual identifies a set of polymorphic forms that distinguishes the individual. See generally National Research Council, The Evaluation of Forensic DNA Evidence (Eds. Pollard et al., National Academy Press, DC, 1996). The more sites that are analyzed, the lower the probability that the set of polymorphic forms in one individual is the same as that in an unrelated individual. Preferably, if multiple sites are analyzed, the sites are unlinked. Thus, polymorphisms of the invention are often used in conjunction with polymorphisms in distal genes. Preferred polymorphisms for use in forensics are biallelic because the population frequencies of two polymorphic forms can usually be determined with greater accuracy than those of multiple polymorphic forms at multi-allelic loci.

The capacity to identify a distinguishing or unique set of forensic markers in an individual is useful for forensic analysis. For example, one can determine whether a blood sample from a suspect matches a blood or other tissue sample from a crime scene by determining whether the set of polymorphic forms occupying selected polymorphic sites is the same in the suspect and the sample. If the set of polymorphic markers does not match between a suspect and a sample, it can be concluded (barring experimental error) that the suspect was not the source of the sample. If the set of markers does match, one can conclude that the DNA from the suspect is consistent with that found at the crime scene. If frequencies of the polymorphic forms at the loci tested have been determined (e.g., by analysis of a suitable population of individuals),

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one can perform a statistical analysis to determine the probability that a match of suspect and crime scene sample would occur by chance.

p(ID) is the probability that two random individuals have the same polymorphic or allelic form at a given polymorphic site. In biallelic loci, four 5 genotypes are possible: AA, AB, BA, and BB. If alleles A and B occur in a haploid genome of the organism with frequencies x and y, the probability of each genotype in a diploid organism is (see WO 95/12607):

Homozygote: $p(AA)=x^2$ Homozygote: $p(BB)=y^2=(1-x)^2$ Single Heterozygote: p(AB)=p(BA)=xy=x(1-x)

Both Heterozygotes: p(AB+BA)= 2xy = 2x(1-x)

The probability of identity at one locus (i.e, the probability that two individuals, picked at random from a population will have identical polymorphic forms at a given locus) is given by the equation:

15 p(ID) =
$$(x^2)^2 + (2xy)^2 + (y^2)^2$$
.

These calculations can be extended for any number of polymorphic forms at a given locus. For example, the probability of identity p(ID) for a 3-allele system where the alleles have the frequencies in the population of x, y and z, respectively, is equal to the sum of the squares of the genotype frequencies:

20
$$p(ID) = x^4 + (2xy)^2 + (2yz)^2 + (2xz)^2 + z^4 + y^4$$

In a locus of n alleles, the appropriate binomial expansion is used to calculate p(ID) and p(exc).

The cumulative probability of identity (cum p(ID)) for each of multiple unlinked loci is determined by multiplying the probabilities provided by each locus.

25 cum
$$p(ID) = p(ID1)p(ID2)p(ID3).... p(IDn)$$

The cumulative probability of non-identity for n loci (i.e. the probability that two random individuals will be different at 1 or more loci) is given by the equation:

cum p(nonID) = 1-cum p(ID).

If several polymorphic loci are tested, the cumulative probability of non-30 identity for random individuals becomes very high (e.g., one billion to one). Such probabilities can be taken into account together with other evidence in determining the guilt or innocence of the suspect.

B. Paternity Testing

The object of paternity testing is usually to determine whether a male is the father of a child. In most cases, the mother of the child is known and thus, the mother's contribution to the child's genotype can be traced. Paternity testing 5 investigates whether the part of the child's genotype not attributable to the mother is consistent with that of the putative father. Paternity testing can be performed by analyzing sets of polymorphisms in the putative father and the child.

If the set of polymorphisms in the child attributable to the father does not match the set of polymorphisms of the putative father, it can be concluded, barring experimental error, that the putative father is not the real father. If the set of polymorphisms in the child attributable to the father does match the set of polymorphisms of the putative father, a statistical calculation can be performed to determine the probability of coincidental match.

The probability of parentage exclusion (representing the probability that a random male will have a polymorphic form at a given polymorphic site that makes him incompatible as the father) is given by the equation (see WO 95/12607):

$$p(exc) = xy(l-xy)$$

where x and y are the population frequencies of alleles A and B of a biallelic polymorphic site.

(At a triallelic site p(exc) = xy(1-xy) + yz(1-yz) + xz(1-xz) + 3xyz(1-xyz)), where x, y and z and the respective population frequencies of alleles A, B and C).

The probability of non-exclusion is

p(non-exc) = 1-p(exc)

The cumulative probability of non-exclusion (representing the value obtained 25 when n loci are used) is thus:

cum p(non-exc) = p(non-exc1)p(non-exc2)p(non-exc3).... p(non-excn)

The cumulative probability of exclusion for n loci (representing the probability that a random male will be excluded)

cum p(exc) = 1 - cum p(non-exc).

If several polymorphic loci are included in the analysis, the cumulative probability of exclusion of a random male is very high. This probability can be taken into account in assessing the liability of a putative father whose polymorphic marker set matches the child's polymorphic marker set attributable to his/her father.

C. Correlation of Polymorphisms with Phenotypic Traits

The polymorphisms of the invention may contribute to the phenotype of an organism in different ways. Some polymorphisms occur within a protein coding

-16-

sequence and contribute to phenotype by affecting protein structure. The effect may be neutral, beneficial or detrimental, or both beneficial and detrimental, depending on the circumstances. For example, a heterozygous sickle cell mutation confers resistance to malaria, but a homozygous sickle cell mutation is usually lethal. Other polymorphisms occur in noncoding regions but may exert phenotypic effects indirectly via influence on replication, transcription, and translation. A single polymorphism may affect more than one phenotypic trait. Likewise, a single phenotypic trait may be affected by polymorphisms in different genes. Further, some polymorphisms predispose an individual to a distinct mutation that is causally related to a certain phenotype.

Phenotypic traits include diseases that have known but hitherto unmapped genetic components (e.g., agammaglobulimenia, diabetes insipidus, Lesch-Nyhan syndrome, muscular dystrophy, Wiskott-Aldrich syndrome, Fabry's disease, familial hypercholesterolemia, polycystic kidney disease, hereditary spherocytosis, von 15 Willebrand's disease, tuberous sclerosis, hereditary hemorrhagic telangiectasia, familial colonic polyposis, Ehlers-Danlos syndrome, osteogenesis imperfecta, and acute intermittent porphyria). Phenotypic traits also include symptoms of, or susceptibility to, multifactorial diseases of which a component is or may be genetic, such as autoimmune diseases, inflammation, cancer, diseases of the nervous system, 20 and infection by pathogenic microorganisms. Some examples of autoimmune diseases include rheumatoid arthritis, multiple sclerosis, diabetes (insulin-dependent and non-independent), systemic lupus erythematosus and Graves disease. Some examples of cancers include cancers of the bladder, brain, breast, colon, esophagus, kidney, leukemia, liver, lung, oral cavity, ovary, pancreas, prostate, skin, stomach and 25 uterus. Phenotypic traits also include characteristics such as longevity, appearance (e.g., baldness, obesity), strength, speed, endurance, fertility, and susceptibility or receptivity to particular drugs or therapeutic treatments.

The correlation of one or more polymorphisms with phenotypic traits can be facilitated by knowledge of the gene product of the wild type (reference) gene. The genes in which cSNPs of the present invention have been identified are genes which have been previously sequenced and characterized in one of their allelic forms. For example, genes of the present invention in which cSNPs have been identified include genes encoding antithrombin III (Humphries, Semin Hematol 32:8-16 (1995); Mammen, Semin Hematol 32:2-6 (1995)), cholesterol ester transfer protein (Bruce and Tall, Curr Opin Lipidol 6:306-311 (1995)), CLanalog (HDL/scavenger receptor) (Freeman, Curr Opin Hematol 4:41-47 (1997); Knecht and Glass, Adv Genet 32:141-198 (1995); Rigotti et al., Curr Opin Lipidol 8:181-188 (1997)), thrombin receptor

(Brass and Molino, Thromb Haemost 78:234-241 (1997); Jamieson, Thromb Haemost 78:242-246 (1997)), thrombin (Eisenberg, Coron Artery Dis 7:400-408 (1996); Jamieson, Thromb Haemost 78:242-246 (1997)), and heparin cofactor II (Bick and Pegram, Semin Thromb Hemost 20:109-132 (1994)). Also included are the genes 5 encoding HMG coA-reductase (Bjelajac et al., Ann Pharmacother 30:1304-1315 (1996)), platelet glycoprotein IIB and IIIA (Jamieson, Thromb Haemost 78:242-246 (1997); Lefkovits et al., N Engl J Med 332:1553-1559 (1995); Nurden, Thromb Haemost 74:345-351 (1995)), lecithin:cholesterol acyltransferase (Kuivenhoven et al., J Lipid Res 38:191-205 (1997)), LDL receptor (Holvoet and Collen, Curr Opin 10 Lipidol 8:320-328 (1997); Rigotti et al., Curr Opin Lipidol 8:181-188 (1997)). protein C (Bertina, Clin Chem 43:1678-1683 (1997); Bick and Pegram, Semin Thromb Hemost 20:109-132 (1994); Humphries, Semin Hematol 32:8-16 (1995); Koeleman et al., Semin Hematol 34:256-264 (1997)), platelet activating factor receptor (Feuerstein et al., J Lipid Mediat Cell Signal 15:255-284 (1997); Shimizu 15 and Mutoh, Adv Exp Med Biol 407:197-204 (1997)), tissue factor (Abildgaard, Blood Coagul Fibrinolysis 6:S45-49(1995); Bick and Pegram, Semin Thromb Hemost 20:109-132 (1994); Harker et al., Haemostasis 1:76-82 (1996); Ruf and Edgington, Faseb J 8:385-390 (1994)), tissue factor pathway inhibitor (Shimizu and Mutoh, Adv Exp Med Biol 407:197-204 (1997); Feuerstein et al., J Lipid Mediat Cell Signal 20 15:255-284 (1997)), thromboxane A2 receptor (Feuerstein et al., J Lipid Mediat Cell

20 15:255-284 (1997)), thromboxane A2 receptor (Feuerstein et al., J Lipid Mediat Cel Signal 15:255-284 (1997); Kinsella et al., Ann NY Acad Sci 714:270-278 (1994); Patrono and Renda, Am J Cardiol 80:17E-20E (1997)), lipoprotein lipase (Applebaum-Bowden, Curr Opin Lipidol 6:130-135 (1995)), and factor V (Bertina, Clin Chem 43:1678-1683 (1997); Harker et al., Haemostasis 1:76-82 (1996);

25 Koeleman et al., Semin Hematol 34:256-264 (1997)).

Correlation is performed for a population of individuals who have been tested for the presence or absence of a phenotypic trait of interest and for polymorphic markers sets. To perform such analysis, the presence or absence of a set of polymorphisms (i.e. a polymorphic set) is determined for a set of the individuals, some of whom exhibit a particular trait, and some of which exhibit lack of the trait. The alleles of each polymorphism of the set are then reviewed to determine whether the presence or absence of a particular allele is associated with the trait of interest. Correlation can be performed by standard statistical methods such as a k-squared test and statistically significant correlations between polymorphic form(s) and phenotypic characteristics are noted. For example, it might be found that the presence of allele A1 at polymorphism A correlates with heart disease. As a further example, it might

be found that the combined presence of allele A1 at polymorphism A and allele B1 at polymorphism B correlates with increased milk production of a farm animal.

Such correlations can be exploited in several ways. In the case of a strong correlation between a set of one or more polymorphic forms and a disease for which 5 treatment is available, detection of the polymorphic form set in a human or animal patient may justify immediate administration of treatment, or at least the institution of regular monitoring of the patient. Detection of a polymorphic form correlated with serious disease in a couple contemplating a family may also be valuable to the couple in their reproductive decisions. For example, the female partner might elect to 10 undergo in vitro fertilization to avoid the possibility of transmitting such a polymorphism from her husband to her offspring. In the case of a weaker, but still statistically significant correlation between a polymorphic set and human disease. immediate therapeutic intervention or monitoring may not be justified. Nevertheless. the patient can be motivated to begin simple life-style changes (e.g., diet, exercise) 15 that can be accomplished at little cost to the patient but confer potential benefits in reducing the risk of conditions to which the patient may have increased susceptibility by virtue of variant alleles. Identification of a polymorphic set in a patient correlated with enhanced receptiveness to one of several treatment regimes for a disease indicates that this treatment regime should be followed.

For animals and plants, correlations between characteristics and phenotype are useful for breeding for desired characteristics. For example, Beitz et al., US 5,292,639 discuss use of bovine mitochondrial polymorphisms in a breeding program to improve milk production in cows. To evaluate the effect of mtDNA D-loop sequence polymorphism on milk production, each cow was assigned a value of 1 if variant or 0 if wildtype with respect to a prototypical mitochondrial DNA sequence at each of 17 locations considered. Each production trait was analyzed individually with the following animal model:

Y_{ijkpn} = μ + YS_i + P_j + X_k + β₁ + ... β₁₇ + PE_n + a_n + e_p
where Y_{ijkmp} is the milk, fat, fat percentage, SNF, SNF percentage, energy
concentration, or lactation energy record; μ is an overall mean; YS_i is the effect common to all cows calving in year-season; X_k is the effect common to cows in either the high or average selection line; β₁ to β₁₇ are the binomial regressions of production record on mtDNA D-loop sequence polymorphisms; PE_n is permanent environmental effect common to all records of cow n; a_n is effect of animal n and is composed of the
additive genetic contribution of sire and dam breeding values and a Mendelian sampling effect; and e_p is a random residual. It was found that eleven of seventeen polymorphisms tested influenced at least one production trait. Bovines having the

best polymorphic forms for milk production at these eleven loci are used as parents for breeding the next generation of the herd.

D. Genetic Mapping of Phenotypic Traits

The previous section concerns identifying correlations between phenotypic traits and polymorphisms that directly or indirectly contribute to those traits. The present section describes identification of a physical linkage between a genetic locus associated with a trait of interest and polymorphic markers that are not associated with the trait, but are in physical proximity with the genetic locus responsible for the trait and co-segregate with it. Such analysis is useful for mapping a genetic locus associated with a phenotypic trait to a chromosomal position, and thereby cloning gene(s) responsible for the trait. See Lander et al., Proc. Natl. Acad. Sci. (USA) 83, 7353-7357 (1986); Lander et al., Proc. Natl. Acad. Sci. (USA) 84, 2363-2367 (1987); Donis-Keller et al., Cell 51, 319-337 (1987); Lander et al., Genetics 121, 185-199 (1989)). Genes localized by linkage can be cloned by a process known as directional cloning. See Wainwright, Med. J. Australia 159, 170-174 (1993); Collins, Nature Genetics 1, 3-6 (1992).

Linkage studies are typically performed on members of a family. Available members of the family are characterized for the presence or absence of a phenotypic trait and for a set of polymorphic markers. The distribution of polymorphic markers in an informative meiosis is then analyzed to determine which polymorphic markers co-segregate with a phenotypic trait. See, e.g., Kerem et al., Science 245, 1073-1080 (1989); Monaco et al., Nature 316, 842 (1985); Yamoka et al., Neurology 40, 222-226 (1990); Rossiter et al., FASEB Journal 5, 21-27 (1991).

Linkage is analyzed by calculation of LOD (log of the odds) values. A lod
25 value is the relative likelihood of obtaining observed segregation data for a marker
and a genetic locus when the two are located at a recombination fraction θ, versus the
situation in which the two are not linked, and thus segregating independently
(Thompson & Thompson, Genetics in Medicine (5th ed, W.B. Saunders Company,
Philadelphia, 1991); Strachan, "Mapping the human genome" in The Human Genome
30 (BIOS Scientific Publishers Ltd, Oxford), Chapter 4). A series of likelihood ratios are
calculated at various recombination fractions (θ), ranging from θ = 0.0 (coincident
loci) to θ = 0.50 (unlinked). Thus, the likelihood at a given value of θ is: probability
of data if loci linked at θ to probability of data if loci unlinked. The computed
likelihoods are usually expressed as the log₁₀ of this ratio (i.e., a lod score). For
35 example, a lod score of 3 indicates 1000:1 odds against an apparent observed linkage
being a coincidence. The use of logarithms allows data collected from different

families to be combined by simple addition. Computer programs are available for the calculation of lod scores for differing values of θ (e.g., LIPED, MLINK (Lathrop, Proc. Nat. Acad. Sci. (USA) 81, 3443-3446 (1984)). For any particular lod score, a recombination fraction may be determined from mathematical tables. See Smith et al., Mathematical tables for research workers in human genetics (Churchill, London, 1961); Smith, Ann. Hum. Genet. 32, 127-150 (1968). The value of θ at which the lod score is the highest is considered to be the best estimate of the recombination fraction.

Positive lod score values suggest that the two loci are linked, whereas negative values suggest that linkage is less likely (at that value of θ) than the possibility that the two loci are unlinked. By convention, a combined lod score of +3 or greater (equivalent to greater than 1000:1 odds in favor of linkage) is considered definitive evidence that two loci are linked. Similarly, by convention, a negative lod score of -2 or less is taken as definitive evidence against linkage of the two loci being compared. Negative linkage data are useful in excluding a chromosome or a segment thereof from consideration. The search focuses on the remaining non-excluded chromosomal locations.

IV. Modified Polypeptides and Gene Sequences

The invention further provides variant forms of nucleic acids and corresponding proteins. The nucleic acids comprise one of the sequences described in the Table, column 8, in which the polymorphic position is occupied by one of the alternative bases for that position. Some nucleic acids encode full-length variant forms of proteins. Similarly, variant proteins have the prototypical amino acid sequences encoded by nucleic acid sequences shown in the Table, column 8, (read so as to be in-frame with the full-length coding sequence of which it is a component) except at an amino acid encoded by a codon including one of the polymorphic positions shown in the Table. That position is occupied by the amino acid coded by the corresponding codon in any of the alternative forms shown in the Table.

Variant genes can be expressed in an expression vector in which a variant gene is operably linked to a native or other promoter. Usually, the promoter is a eukaryotic promoter for expression in a mammalian cell. The transcription regulation sequences typically include a heterologous promoter and optionally an enhancer which is recognized by the host. The selection of an appropriate promoter, for example trp, lac, phage promoters, glycolytic enzyme promoters and tRNA promoters, depends on the host selected. Commercially available expression vectors can be used. Vectors can include host-recognized replication systems, amplifiable genes, selectable markers, host sequences useful for insertion into the host genome, and the like.

The means of introducing the expression construct into a host cell varies depending upon the particular construction and the target host. Suitable means include fusion, conjugation, transfection, transduction, electroporation or injection, as described in Sambrook, supra. A wide variety of host cells can be employed for expression of the variant gene, both prokaryotic and eukaryotic. Suitable host cells include bacteria such as *E. coli*, yeast, filamentous fungi, insect cells, mammalian cells, typically immortalized, e.g., mouse, CHO, human and monkey cell lines and derivatives thereof. Preferred host cells are able to process the variant gene product to produce an appropriate mature polypeptide. Processing includes glycosylation, 10 ubiquitination, disulfide bond formation, general post-translational modification, and the like. As used herein, "gene product" includes mRNA, peptide and protein products.

The protein may be isolated by conventional means of protein biochemistry and purification to obtain a substantially pure product, i.e., 80, 95 or 99% free of cell component contaminants, as described in Jacoby, Methods in Enzymology Volume 104, Academic Press, New York (1984); Scopes, Protein Purification, Principles and Practice, 2nd Edition, Springer-Verlag, New York (1987); and Deutscher (ed), Guide to Protein Purification, Methods in Enzymology, Vol. 182 (1990). If the protein is secreted, it can be isolated from the supernatant in which the host cell is grown. If not 20 secreted, the protein can be isolated from a lysate of the host cells.

The invention further provides transgenic nonhuman animals capable of expressing an exogenous variant gene and/or having one or both alleles of an endogenous variant gene inactivated. Expression of an exogenous variant gene is usually achieved by operably linking the gene to a promoter and optionally an enhancer, and microinjecting the construct into a zygote. See Hogan et al., "Manipulating the Mouse Embryo, A Laboratory Manual," Cold Spring Harbor Laboratory. Inactivation of endogenous variant genes can be achieved by forming a transgene in which a cloned variant gene is inactivated by insertion of a positive selection marker. See Capecchi, Science 244, 1288-1292 (1989). The transgene is then introduced into an embryonic stem cell, where it undergoes homologous recombination with an endogenous variant gene. Mice and other rodents are preferred animals. Such animals provide useful drug screening systems.

In addition to substantially full-length polypeptides expressed by variant genes, the present invention includes biologically active fragments of the polypeptides, or analogs thereof, including organic molecules which simulate the interactions of the peptides. Biologically active fragments include any portion of the full-length polypeptide which confers a biological function on the variant gene

product, including ligand binding, and antibody binding. Ligand binding includes binding by nucleic acids, proteins or polypeptides, small biologically active molecules, or large cellular structures.

Polyclonal and/or monoclonal antibodies that specifically bind to variant gene products but not to corresponding prototypical gene products are also provided. Antibodies can be made by injecting mice or other animals with the variant gene product or synthetic peptide fragments thereof. Monoclonal antibodies are screened as are described, for example, in Harlow & Lane, Antibodies, A Laboratory Manual, Cold Spring Harbor Press, New York (1988); Goding, Monoclonal antibodies,

10 Principles and Practice (2d ed.) Academic Press, New York (1986). Monoclonal antibodies are tested for specific immunoreactivity with a variant gene product and lack of immunoreactivity to the corresponding prototypical gene product. These antibodies are useful in diagnostic assays for detection of the variant form, or as an active ingredient in a pharmaceutical composition.

15 V. Kits

The invention further provides kits comprising at least one allele-specific oligonucleotide as described above. Often, the kits contain one or more pairs of allele-specific oligonucleotides hybridizing to different forms of a polymorphism. In some kits, the allele-specific oligonucleotides are provided immobilized to a substrate.

- 20 For example, the same substrate can comprise allele-specific oligonucleotide probes for detecting at least 10, 100 or all of the polymorphisms shown in the Table. Optional additional components of the kit include, for example, restriction enzymes, reverse-transcriptase or polymerase, the substrate nucleoside triphosphates, means used to label (for example, an avidin-enzyme conjugate and enzyme substrate and
- 25 chromogen if the label is biotin), and the appropriate buffers for reverse transcription, PCR, or hybridization reactions. Usually, the kit also contains instructions for carrying out the methods.

The following Examples are offered for the purpose of illustrating the present invention and are not to be construed to limit the scope of this invention. The teachings of all references cited herein are hereby incorporated herein by reference.

EXAMPLES

The polymorphisms shown in the Table were identified by resequencing of target sequences from a minimum of 50 unrelated individuals of diverse ethnic and geographic backgrounds by hybridization to probes immobilized to microfabricated

arrays. The strategy and principles for design and use of such arrays are generally described in WO 95/11995.

A typical probe array used in this analysis has two groups of four sets of probes that respectively tile both strands of a reference sequence. A first probe set 5 comprises a plurality of probes exhibiting perfect complementarily with one of the reference sequences. Each probe in the first probe set has an interrogation position that corresponds to a nucleotide in the reference sequence. That is, the interrogation position is aligned with the corresponding nucleotide in the reference sequence, when the probe and reference sequence are aligned to maximize complementarily between 10 the two. For each probe in the first set, there are three corresponding probes from three additional probe sets. Thus, there are four probes corresponding to each nucleotide in the reference sequence. The probes from the three additional probe sets are identical to the corresponding probe from the first probe set except at the interrogation position, which occurs in the same position in each of the four 15 corresponding probes from the four probe sets, and is occupied by a different nucleotide in the four probe sets. In the present analysis, probes were 25 nucleotides long. Arrays tiled for multiple different references sequences were included on the same substrate.

Publicly available sequences for a given gene were assembled into Gap4

20 (http://www.biozentrum.unibas.ch/~biocomp/staden/Overview.html). PCR primers covering each exon were designed using Primer 3 (http://www-genome.wi.mit.edu/cgi-bin/primer/primer3.cgi). Primers were not designed in regions where there were sequence discrepancies between reads. For CLA1, whose genomic sequence is not published, nested primers were designed from the cDNA. For all genes except CLA1, genomic DNA was amplified in at least 50 individuals using 2.5 pmol each primer, 1.5 mM MgCl₂, 100 μM dNTPs, 0.75 μM AmpliTaq GOLD polymerase, and 19 ng DNA in a 15 μl reaction. Reactions were assembled using a PACKARD MultiPROBE robotic pipetting station and then put in MJ 96-well tetrad thermocyclers (96°C for 10 minutes, followed by 35 cycles of 96°C for 30 seconds, 59°C for 2 minutes, and 72°C for 2 minutes). A subset of the PCR assays for each individual were run on 3% NuSieve gels in 0.5X TBE to confirm that the reaction worked.

For CLA1, first strand cDNA was made using the Gibco BRL SuperScript
Preamplification Kit (#18089-011) and following the manufacturers instructions
35 except that 150 ng of random hexamers were used to primer 1 µg of total RNA. The
cDNA was amplified using the outermost primer pairs and the above conditions; 1/20
of the reaction was used as a template for the secondary PCR using the innermost

primers. All RT-PCR products were run on 2% NuSieve gels in 1X TAE to confirm the presence of a product.

For a given DNA, 5 μl (about 50ng) of each PCR or RT-PCR product were pooled (Final volume = 150-200 μl). The products were purified using QiaQuick

5 PCR purification from Qiagen. The samples were eluted once in 35 μl sterile water and 4 μl 10X One-Phor-All buffer (Pharmacia). The pooled samples were digested with 0.2 μ DNaseI (Promega) for 10 minutes at 37°C and then labeled with 0.5 nmols biotin-N6-ddATP and 15 μ Terminal Transferase (GibcoBRL Life Technology) for 60 minutes at 37°C. Both fragmentation and labeling reactions were terminated by incubating the pooled sample for 15 minutes at 100°C.

Low-density DNA chips (Affymetrix,CA) were hybridized following the manufacturer's instructions. Briefly, the hybridization cocktail consisted of 3M TMACl, 10 mM Tris pH 7.8, 0.01% Triton X-100, 100 mg/ml herring sperm DNA (Gibco BRL), 200 pM control biotin-labeled oligo. The processed PCR products were denatured for 7 minutes at 100°C and then added to prewarmed (37°C) hybridization solution. The chips were hybridized overnight at 44°C. Chips were washed in 1X SSPET and 6X SSPET followed by staining with 2 µg/ml SARPE and 0.5 mg/ml acetylated BSA in 200 µl of 6X SSPET for 8 minutes at room temperature. Chips were scanned using a Molecular Dynamics scanner.

- Chip image files were analyzed using Ulysses (Affymetrix, CA) which uses four algorithms to identify potential polymorphisms. Candidate polymorphisms were visually inspected and assigned a confidence value: high confidence candidates displayed all three genotypes, while likely candidates showed only two genotypes (homozygous for reference sequence and heterozygous for reference and variant).
- 25 Some of the candidate polymoprhisms were confirmed by ABI sequencing. Identified polymorphisms were compared to SwissProt and the Mutation Database to determine if they were novel. Results are shown in the Table.

In the Table, the genes listed in column 2 are as follows: antithrombin III (AT3); cholesterol ester transfer protein (CETP); CLanalog (HDL/scavenger receptor) (CLanalog); thrombin receptor (F2R); thrombin (F2); heparin Cofactor II (HCF2); HMG coA-reductase (HMGCR); platelet glycoprotein IIB (ITGA2B); platelet glycoprotein IIIA (ITGB3); lecithin:cholesterol acyltransferase (LCAT); LDL receptor (LDLR); protein C (PROC); platelet activating factor receptor (PTAFR); tissue factor pathway inhibitor (TFPI); thromboxane A2 receptor (TBXA2R); lipoprotein lipase (LPL); tissue factor (F3); and factor V (F5).

Column 1 of the Table shows the laboratory name for the particular gene.

Column 3 shows the GenBank Accession number for the wild type (reference) allele.

Column 4 shows the nucleotide number location of the polymorphism relative to the numbering of the sequence deposited with GenBank having the listed Accession number; the GenBank sequence is understood to be the nucleotide sequence present in the GenBank database on April 1, 1998, which sequences are incorporated herein by reference in their entirety. These GenBank sequences are illustrated in Figures 3-38.

Column 5 shows the codon which is altered by the polymorphism. Columns 6, 7 and 8 show the reference codon, variant codon and amino acid change, respectively, for the silent polymorphisms. Columns 9, 10 and 11 show the reference codon, variant codon and amino acid change, respectively, for the missense polymorphisms. Columns 12, 13 and 14 show the reference codon, variant codon and amino acid change, respectively, for the nonsense polymorphisms. Columns 15 and 16 show the nucleotide of the reference allele and the frequency of that allele, respectively. This base is arbitrarily designated the reference or prototypical form, but it is not necessarily the most frequently occurring form. Columns 17 and 18 show the nucleotide of the variant allele and the frequency of that allele, respectively. It is noted that the genes with polymorphism IDs of F5u8, HCF2u1 and HMGCRu2 contained the indicated polymorphism at the indicated nucleotide position, but that these nucleotide positions are in the non-coding region of the gene.

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						Silent PM	Æ.	Σ	Missense	e PM	Ż	Nonsense	¥.		Allele Freg.	Je J.	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	Var codon	УУ сряхаде	Ref codon	Var codon	сралде УУ	Ref codon	Var codon	95пълс Брабар	Ref allele	Preq	Var allele	Freq
AT3u3	AT3	M21645	100	438				AGG	GGG	R to G				4	66.0	b	0.01
CETPul	CETP	M30185	1298	390				သည	ccc	A to P				ບ	0.95	υ	0.05
CETPu8	CETP	J02898	298	455				GTG	ATG	V to M				ď	66.0	4	10.0
CETPu9	CETP	302898	125	486		-		GTG	ATG	V to M				b	0.99	4	0.01
CLanalogu3	CLanalog	222555	400	111				GTG	ATG	V to M				ຍ	0.99	4	0.01
CLanalogu4	CLanalog	222555	472	135				GTC	ATC	V to I				b	0.99	4	0.01
F2Ru1	F2R	M62424	496	91				GAT	GGT	D to G				٧	0.99	b	0.01
F2Ru2	F2R	M62424	610	129				DE C	ട്ടാ	L to R				H	96.0	g	0.02
F2Ru3	F2R	M62424	664	147				GCA	GAA	A to E				υ	0.91	K	60.0
F2Ru4	F2R	M62424	720	166				AGT	GGT	S to G				Ą	0.99	_S	10.0
F2Ru6	F2R	M62424	405	61				AAA	CAA	K to Q				4	0.93	υ	0.07
P2u1	F2	M17262	10777	165				ACG	ATG	T to M				υ	0.97	1	0.03
Fzuz	F2	M17262	15342	386				ည	ACC	P to T				υ	0.99	A A	0.01
F3u1	F3	J02846	9363	163 .				gg	TGG	R to W				υ	66.0	į.	0.01
F5u4	FS	M14335	1314	413				ATG	ACG	M to T				н	0.94	υ	90.0
HCF2u3	HCF2	M12849	1353	442				ACG	ATG	T to M				υ	66.0	Ţ	0.01

						Silent	Md .	Σ	Missense	e PM	Ż	Nonsense	MG a		Allele Freg.	le I.	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	Var codon	сучиде УУ	Ref codon	Var codon	AA change	Ref codon	Var codon	AA egnsdə	Kef allele	Freq	Var allele	Fred
HCF2u4	HCF2	M12849	47	7				క్ర	ACA	A to T				υ	0.98	4	0.03
HCF2u6	HCF2	M12849	651	208				ည္တ	CAC	R to H				U	0.99	4	0.01
HMGCRu1	HMGCR	M11058	1962	638				ATA	GTA	I to V				∢	0.99	U	0.01
ITGA2Bu2	ITGA2B	J02764	2623	874				ATC	AGC	I to S			•	H	67.0	5	0.21
ITGA2Bu5	ITGAZB	302764	2904	896				TAT	AAT	Y to N				ь	0.99	4	0.01
ITGA2Bu6	1TGA2B	J02764	120	40				ACC	ATC	T to I				U	0.97	ь	0.03
ITGA2Bu7	ITGAZB	J02764	2299	766				ATT	AGT	I to S				H	66.0		0.01
ITGB3u1	ITGB3	302703	526	169				CGA	ð	R to Q				o	0.99	A	0.01
ITGB3u8	ITGB3	502703	1377	453				GTC	ATC	V to I	·			ט	66.0	4	0.01
LCATu2	LCAT	M12625	196	232				TCT	ACT.	S to T				F+	96.0	4	0.02
LDLRu14	LDLR	L00351	69	814				ဗ္ဗ	S S	R to Q				ט	0.99	4	0.01
LDLRu7	LDLR	L29401	169	2				999	990	G to R				U	0.99	U	0.01
LDLRu8	LDLR	L00344	59	468				GTC	ATC	V to I				Ö	66.0	4	0.01
2n า ๔า	Tat	M15856	1453	427				229	ACC	A to T				U	0.99	4	0.01
PROCu4	PROC	K02059	534	283				AAG	AGG	K to R				4	66.0	,	0.01
PTAFRu3	PTAFR	D10202	783	224				ţj	GAT	A to D				υ	0.99	4	0.01
PTAFRU4	PTAFR	D10202	194	28				CTC	TTC	L to F				υ	0.99	F	0.01

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						Silent PM	. PM	ž	Missense	e PM	z	Nonsense	W. as		Allele Freq.] ie	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	ляк содоп	AA change	Ref codon	Var codon	УУ сучиде	Ref codon	Var codon	сруиде УУ	Ref allele	Fred	Var allele	Freq
PTAFRUS	PTAFR	D10202	1125	338				AAT	AGT	N to S				4	86.0	b	0.02
TPPIul	TFPI	J03225	1006	292				GTG	ATG	V to M				ט	66.0	4	0.01
CETPu4	CETP	M30185	196	22	ACC	ACA	T to T							υ	66.0	4	0.01
LDLRu13	LDLR	100336	29	7.2	TGT	TGC	C to C							T	0.62	U	0.38
HCP2u2	нсга	M12849	259	77	GAC	GAT	D to D							၁	. 76.0	1	0.03
CETPu5	CETP	M30185	388	. 86	ATC	ATT	I to I							υ) 66.0	o o	0.01
HCF2u5	HCP2	M12849	313	95	ATC	ATT	I to I							၁	0.99	£	0.01
1TGB3u7	ITGB3	502703	362	114	ATT	ATC	I to I							7	0.97	υ	0.03
F2Ru7	P2R	M62424	609	129	CTG	TTG	L to L							υ	1 86.0	٠ 1	0.02
PROCu2	PROC	K02059	109	141	TCT	TCG	S to S							T	0.46	0	0.54
CLanalogu2	CLanalog	222555	570	167	၁၅၅	GGT	G to G							υ	0.88	٠ ۲	0.12
F2RuS	F2R	M62424	740	172	TCT	TCG	S to S							Т	0.99	U	0.01
LCATUI	LCAT	M12625	864	199	GTC	GTT	V to V							υ	0.99 T	-	0.01,
CETPu6	CETP	M30185	766	212	၁၁၁	GCT	A to A							υ	0.98 T		0.02
PROCu3	PROC	M11228	9358	256	GAT	GAC	D to D							Ţ	0.98 C	0	.02
F2u4	F2	M17262	13434	271	ပ္ပပ္ပ	GGT	G to G					-		υ	1 86.0	0	.02
ITGB3u3	ITGB3	302703	902	294	t)	သသ	P to P						-	Ę+	0.87 C		0.13

						Silent PM	W J	Ξ	Missense	e PM	z	Nonsense	₩.		Allele Freq.] ie	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	Var codon	epange change	Ref codon	Var codon	УУ срадде	Ref codon	Var codon	сралде УУ	Ref allele	Ered	Var allele	Freq
PROCu1	PROC	K02059	577	297	GAC	GAT	D to D							υ	66.0	н	0.01
LCATu4	LCAT	M12625	1167	300	œr	၁၅၁	R to R							н	0.99	υ	0.01
CLanalogu5	CLanalog	222555	972	301	TTC	TT	F to F							υ	0.95	£-	0.05
TBXA2Ru1	TBXA2R	D38081	1915	308	TAT	TAC	Y to Y							Ŧ	0.57	Ü	0.43
AT3u1	AT3	D29832	1005	327	GTG	GTA	V to V							ຽ	0.64	٧	0.36
CLanalogul	CLanalog	22255	1119	350	ວວວ	GCT	A to A							၁	0.68	ŀ	0.32
ITGB3u4	ITGB3	302703	1163	381	GTC	GTA	V to V							υ	0.50	۷.	0.50
LPLu1	LPL	M15856	1338	388	ACC	ACA	T to T							υ	0.89	4	0.11
LCATu3	LCAT	M12625	1444	393	CT SEC	TTG	L to L							υ	0.93	F .	0.07
F2u3	F2	M17262	15419	411	ວວວ	CC.	P to P							ť	76.0	4	0.03
Psus	FS	M14335	1318	414	AAA	AAG	К to К							4	0.92	ט	90.0
CETPu7	CETP	M30185	1429	433	GTG	GTA	V to V							ŋ	, 66.0	4	0.01
LDLRu9	LDLR	L00343	152	441.	ATC	ATT	I to I							υ	. 66.0	4	0.01.
AT3u4	AT3	D29832	1374	450	AAC	AAT	N to N							υ	0.99	4	0.01
PSul	PS	M14335	1456	460	AAC	AAT	N to N							ပ	0.95	T	0.05
HCF2u7	HCF2	M12849	1474	482	CAC	CAT	н со н							Ü	0.53	٠	0.47
ITGB3u5	ITGB3	M20311	1549	511	GAG	g S	路 to 路							ט	0.27	4	0.73

						Silent	Md.	Σ	Missense PM	Ψ. W.	ž	Nonsense	PW e		Allele Freq.	e .	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	Var codon	у сучиде Сучи	Ref codon	Var codon	сузиде УУ	Ref codon	Var codon	срэпде У У	Ref allele	Freq	Var allele	Freq
ITGB3u6	ITGB3	M20311	1561	515	CGA	වසුර	R to R							æ	0.43	9	0.57
F2u5	23	M17262	16827	534	CCC	CCA	P to P							υ	66.0	4	0.01
LDLRu3	LDLR	L00345	46	539	CCC	ccr	P to P							υ	0.89	F	11.0
PSu6	PS	M14335	1792	572	GAG	GAA	E to E		-					ט	0.94	A	90.0
LDLRu10	מדמיז	059436	45	575	crc	crr	L to L							υ	0.93	[+	0.07
LDLRu6	מתמד	U59436	93	591	AAT	AAC	N to N							Ę-i	0.77	U	0.23
ITGAZBu3	ITGA2B	M33320	6845	605	ഡ	භි	P to P							ט	0.98	4	0.02
LDLRu11	LDLR	L00347	90	640	AAC	AAT	N to N							υ	0.99	ь	0.01
F5u7	FS	M14335	2002	642	ACC	ş	T to T							υ	96.0	<u> </u>	0.04
LDf.Ru1	LDLR	L00347	129	653	GTC	GTT	V to V							υ	0.31	F	69.0
LDLRu12	LDLR	L00349	107	744	ccc	CGA	R to R							Ü	0.85	٠,	0.15
ITGA2Bu8	ITGA2B	J02764	2567	855	crr	CTC	L to L							E٠	0.99	υ	0.01
ITGA2Bu4	ITGA2B	302764	2918	972	ട്ടാ	ģ	P to P							υ	66.0	4	0.01,
ITGA2Bu1	ITGA2B	M22569	194	1021	GTC	GTT	V to V							υ	0.66	ь	0.34
P5u8	F5	L32765	66											ט	0.99	ь	0.01
HCP2u1	HCF2	M58600	11907											Ü	0.96	F	0.04
HMGCRu2	HMGCR	M11058	2725						\neg					5	0.97 A		0.03

						Silent PM	. PM	Σ	Missense PM	e PM	z	Nonsense PM	e PM		All Fre	Allele Freq.	
Polymorphism ID	Gene	GenBank Accession No.	Nt. Position	Codon No.	Ref codon	Var codon	еривиде Суч	Ref codon	иороэ тьу	сряиде УУ	Ref codon	Var codon	уу сруиде	Ref allele	Freq	Var allele	Freq
ITGB3u2	ITGB3	502703	961	59				ന്ദ	ອວວ	L to P				Ļ	0.87	υ	0.13
CETPu2	CETP	M30185	1394	422.				ATC	GTC	I to V				4	0.34	g	99.0
F5u2	F5	M14335	1614	513				AGA	AAA	R to K				ŋ	0.85	Ą	0.15
PSu3	PS	M14335	1677	534				CGA	CAA	R to Q				ย	0.99	4	0.01
AT3u2	AT3	D29832	1035	337	CAG	CAA	0 to 0							U	0.62	4	0.38
LDLRuS	LDLR	L00344	70	471	AGG	AGA	R to R							υ	0.68	4	0.32
LPLu3	747	M76722	3150	474							ភ្ជ	Į.	S to *	υ	0.85	U	0.15

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Genotyping and genetic association studies were performed with respect to the allelic forms of the F5U4 and HCF2U4 genes, and the presence of the reference and variant alleles (as shown in Table 1) were correlated with the occurrence of venous thrombosis and pulmonary emboli. The results are shown in Tables 2 and 3.

TABLE 2: HCF2U4 GENETIC ASSOCIATION STUDY

	Case	Control
Reference	115	115
Heterozygote	5	0

(p = 0.027 by Chi-square test)

(p = 0.06 by Fisher's exact test (two-tailed)).

The F5u4 variant leads to an amino acid substitution (Met413Thr) in the coagulation factor V gene. Another common variant in Factor V (Arg506Gln), the Leiden Variant, is the most common genetic factor predisposing to thrombosis that has been identified to date. Genotyping of patients with deep venous thrombosis has confirmed a statistical association of this variant with deep venous

thrombosis/pulmonary embolism in two separate populations of patients, as shown below:

TABLE 3: F5U4 GENETIC ASSOCIATION STUDY

	REF	HET	VAR	ТОТАІ	ALLELI	E FREQ
	KLI	nei	VAR	TOTAL	REF	VAR
Case	226	38	5	269	91%	9%
Control	207	28	0	235	94%	6%

20 2nd Population

Case	85	28	2	115	86%	14%
Control	95	14	4	113	90%	10%

(p < 0.05 by Chi-square test for combined populations)

These data indicate that there is a trend toward an association between the presence of the variant allele (or heterozygousity) and the occurrence of venous thrombosis and/or pulmonary emboli.

From the foregoing, it is apparent that the invention includes a number of general uses that can be expressed concisely as follows. The invention provides for the use of any of the nucleic acid segments described above in the diagnosis or monitoring of diseases, such as cancer, inflammation, heart disease, diseases of the cardiovascular system, and infection by microorganisms. The invention further provides for the use of any of the nucleic acid segments in the manufacture of a medicament for the treatment or prophylaxis of such diseases. The invention further provides for the use of any of the DNA segments as a pharmaceutical.

All references cited above are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication or patent application were specifically and individually indicated to be so incorporated by reference.

While this invention has been particularly shown and described with references to preferred embodiments thereof, it will be understood by those skilled in the art that various changes in form and details may be made therein without departing from the spirit and scope of the invention as defined by the appended claims.

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CLAIMS

WE CLAIM:

- 1. A nucleic acid molecule selected from the group consisting of the genes listed in the Table, wherein said nucleic acid molecule is at least 5 nucleotides in length and comprises a polymorphic site identified in the Table, wherein a nucleotide at the polymorphic site is different from a nucleotide at the polymorphic site in a corresponding reference allele.
 - 2. A nucleic acid molecule according to Claim 1, wherein said nucleic acid molecule is at least 10 nucleotides in length.
- 10 3. A nucleic acid molecule according to Claim 1, wherein said nucleic acid molecule is at least 20 nucleotides in length.
 - 4. A nucleic acid molecule according to Claim 1, wherein the nucleotide at the polymorphic site is the variant nucleotide for the gene listed in the Table.
- 5. An allele-specific oligonucleotide that hybridizes to a portion of a gene selected from the group consisting of the genes listed in the Table, wherein said portion is at least 5 nucleotides in length and comprises a polymorphic site identified in the Table, wherein a nucleotide at the polymorphic site is different from a nucleotide at the polymorphic site in a corresponding reference allele.
- 20 6. An allele-specific oligonucleotide according to Claim 5 that is a probe.
 - 7. An allele-specific oligonucleotide according to Claim 5, wherein a central position of the probe aligns with the polymorphic site of the portion.
 - 8. An allele-specific oligonucleotide according to Claim 5 that is a primer.
- An allele-specific oligonucleotide according to Claim 8, wherein the 3' end of
 the primer aligns with the polymorphic site of the portion.

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- 10. An isolated gene product encoded by a nucleic acid molecule according to Claim 1.
- 11. A method of analyzing a nucleic acid sample, comprising obtaining the nucleic acid from an individual sample; and determining a base occupying any one of the polymorphic sites shown in the Table.
- 12. A method according to Claim 11, wherein the nucleic acid sample is obtained from a plurality of individuals, and a base occupying one of the polymorphic positions is determined in each of the individuals, and the method further comprising testing each individual for the presence of a disease phenotype, and correlating the presence of the disease phenotype with the base.

				. &	SILENT POLYMORPHISMS	T HISMS	Ю	MISSENSE POLYMORPHISMS	ISE HISMS	&	NONSENSE POLYMORPHISMS	SE		ALLELE PREQUENCIES	CIES	
Polymorphism ID	Gene		Codon No.	Ref codon	Var codon	уу Сувиде УУ	Ref codon	лят содоп	суулде УУ	Ref codon	Var codon	УУ сралуе А	Ref allele	Freq	Var allele	Freq
AT3u3	AT3	•	438				AGG	999	R to G				٧	0.99	U	0.01
CETPu1	CETP		390				၁၁၅	သသ	A to P				υ	0.95	o o	0.05
CETPu8	CETP		455				GTG	ATG	V to M				g	0.99	4	0.01
CETPu9	CETP		486				GTG	ATG	V to M				0	66.0	A 0	0.01
CLanalogu3	CLanalog		111				GTG	ATG	V to M				0	66.0	٧	0.01
CLanalogu4	CLanalog		135				GTC	ATC	V to I				Ö	0.99	0	0.01
F2Ru1	F2R		91				GAT	GGT	D to G				٧	0.99	0	0.01
F2Ru2	F2R		129				CTG	ടാ	L to R				Ð	96.0	0	0.02
F2Ru3	F2R		147				GCA	GAA	A to E				၁	0.91	4	0.09
P2Ru4	P2R		166				AGT	GGT	S to G				4	0.99	0	0.01
F2Ru6	F2R	·	61				AAA	CAA	K to Q				٧	0.93	υ	0.07
F2u1	P2		165				ACG	ATG	T to M				၁	T 76.0		0.03
F2u2	P2		386				သသ	ACC	P to T				၁	66.0	٨ 0	0.01
F3u1	P3		163				ອອວ	TGG	R to W				υ	0.99 T	_	0.01
FSud	PS		413				ATG	ACG	M to T				Į.	0.94 C		90.0
HCP2u3	HCP2		442				ACG	ATG	T to M				υ	0.99 T		0.01
HCP2u4	HCF2		7				gC _A	ACA	A to T				O	0.98 A		0.02
HCP2u6	HCF2		208				၁၅၁	CAC	R to H				О	A 66.0		0.01

FIG. 1A

					24	SILENT POLYMORPHISMS	NT PHISMS	2	MISSENSE POLYMORPHISMS	ISE HISMS	8	NONSENSE POLYMORPHISMS	SE		ALLELE PREQUENCIES	CIES	
Polymorphism ID	Gene			Codon No.	Ref codon	Var codon	су з иде УУ	Ref codon	Var codon	cysude YY	Ref codon	Var codon	суулде УУ	Ref allele	Freq	Var allele	Freq
HMGCRu1	HMGCR			638				ATA	GTA	I to V				0	0.99	0	0.01
ITGA2Bu2	ITGAZB			874				ATC	AGC	I to S				٠ ۲	0.79	0	.21
ITGA2Bu5	ITGA2B			968				TAT	AAT	Y to N				т 0.	66	0 4	0.01
ITGA2Bu6	ITGA2B			40		,		ACC	ATC	T to I				0 0	0.97	1 0	0.03
ITGA2Bu7	ITGA2B			766				ATT	AGT	I to S				٠ ٢	0.99	0 0	0.01
ITGB3u1	1TGB3			169				CGA	CAS	R to Q				0 0	0.99 A	-	0.01
ITGB3u8	ITGB3			453				GTC	ATC	V to I				G 0	0.99 A		0.01
LCATU2	LCAT			232				TCT	ACT	S to T				т 0.	A 86.		0.02
LDLRu14	นาดา			814				ອອວ	CAG	R to Q				G 0	A 66		0.01
LDLRu7	นาตา			2				GGG	ടടാ	G to R				0	0.99 c		0.01
LDLRu8	בטנת			468				GTC	ATC	V to I			,	0 5	0.99 A		0.01
LPLu2	าสา	-		427				၁၁၅	ACC	A to T			,	G 0	0.99 A	-	0.01
PROCu4	PROC			283				AAG	AGG	K to R				٥ ٨	0.99 G		0.01
PTAPRu3	PTAPR		·	224				GCT	GAT	A to D			J	0 0	0.99 A	H	0.01
PTAFRu4	PTAFR			28				CTC	TTC	L to F			Ì	o U	0.99 T	-	0.01
PTAFRUS	PTAFR			338				AAT	AGT	N to S				0	0.98 G		0.02
TPPIU1	IGAL			292				GTG	ATG	V to M			,	0	0.99 A		0.01
CBTPu4	CETP			22	ACC	ACA	T to T						,	٥	0.99 A		0.01
LDLRu13	נסנת			27	ŢĢ.	TGC	C to C							٠ ا	0.62 C		0.38

FIG. 1B

				.	SILENT POLYMORPHISMS	NT PHISMS	TOA	MISSENSE POLYMORPHISMS	SE	Ĭ.	NONSENSE POLYMORPHISMS	SE		ALLELE PREQUENCIES	ILE ACIES	
Polymorphism ID	Gene		Codon No.	Ref codon	Var codon	суучде УУ	Ref codon	Var codon	среиде УУ	Ref codon	Var codon	cysude YY	Ref allele	Freq	Var allele	Freq
HCF2u2	HCF2		7.7	GAC	GAT	D to D							b	0.97	T	0.03
CETPu5	CETP		86	ATC	ATT	I to I							υ	0.99	Ŀ	0.01
HCF2u5	HCF2		95	ATC	ATT	I to I							ပ	0.99	F-	0.01
ITGB3u7	ITGB3		114	ATT	ATC	I to I							Ŧ.	0.97	υ	0.03
F2Ru7	F2R		129	CTG	TTG	L to L							၁	96.0	į.	0.03
PROCu2	PROC		141	TCT	TCG	S to S							Ŧ	0.46	0	0.54
CLanalogu2	CLanalog		167	299	GGT	G to G							ر ن	0.88	7	0.12
F2Ru5	F2R		172	TCT	тсв	S to S							7	0.99	0	0.01
LCATu1	LCAT		199	GTC	GTT	V to V							υ	0.99	e E	0.01
CETPu6	CETP		212	၁၁၅	GCT	A to A					·		၁	96.0	1 0	0.02
PROCu3	PROC		256	GAT	GAC	D to D							4	96.0	0	0.02
F2u4	F2		271	၁၅၅	GGT	G to G							٥	96.0	4	0.02
ITGB3u3	ITGB3		294	CCT	သသ	P to P						,	ę,	0.87	υ	0.13
PROCu1	PROC		297	GAC	GAT	D to D							0 0	0.99	T 0	0.01
LCATu4	LCAT		300	CGT	၁၅၁	R to R							T (66.0	0	0.01,
CLanalogu5	CLanalog		301	TTC	TT	P to F						-	0 0	.95	٠ 0	0.05
TBXA2Ru1	TBXA2R		308	TAT	TAC	YtoY							1	0.57	0 0	0.43
AT3u1	AT3		327	GTG	GTA	v to v							v	0.64	٥ م	0.36
CLanalogu1	CLanalog		350	၁၁၅	GCT	A to A							٥	99.0	T 0	0.32

16. J

				X	SILENT POLYMORPHISMS	NT PH I SMS	Ю	MISSENSE POLYMORPHISMS	ISE IISMS	102	NONSENSE POLYMORPHISMS	SE		ALLELE FREQUENCTES	LE ACTES	
Polymorphism ID	Gene	ē	Codon No.	Ref codon	ляк содои	сувиде УУ	Ref codon	Var codon	сууиде УУ	Ref codon	Var codon	сууиде УУ	Ref allele	Ered	Var allele	Freq
ITGB3u4	1TGB3		381	GTC	GTA	V to V							υ	0.50	4	0.50
LPLu1	LPL		388	ACC	ACA	T to T							υ	0.89	4	0.11
LCATu3	LCAT		393	CTG	TTG	L to L							၁	0.93	T	0.07
F2u3	F2		411	၅၁၁	ccA	PtoP							U	76.0	V V	0.03
P5u5	F5		414	AAA	AAG	K to K							٧	0.92	٥	0.08
CETPu7	CETP		433	GTG	GTA	V to V		-					9	0.99	A .	0.01
LDLRu9	LDLR		441	ATC	ATT	I to I							၁	0.99	T (0.01
AT3u4	AT3		450	AAC	AAT	N to N							υ	0.99	T (0.01
F5u1	PS		460	AAC	AAT	N to N							υ	96.0	£-	0.05
HCF2u7	HCF2		482	CAC	CAT	н со н							υ	0.53	T (0.47
ITGB3uS	ITGB3		511	GAA	GAA	E to E							9	0.27	٧	0.73
ITGB3u6	ITGB3		515	ടടാ	ളൊ	R to R							A	0.43	0	0.57
F2u5	F2		534	ಖಂ	CCA	P to P							υ	0.99	A A	0.01
LDLRu3	רטנה		539	၁၁၁	ccr	P to P							υ	0.89	F.	0.11
F5u6	75		572	GAG	GAA	E to E							o o	0.94	٨	0.06
LDLRu10	רטנא		575	CTC	CTT	L to L							υ	0.93	4	0.07
LDLRu6	רטנת		591	AAT	AAC	N to N							į.	77.0	υ υ	0.23
ITGA2Bu3	ITGA2B		605	_D	S S S S	P to P							0	96.0	4	0.02
LDLRu11	LDLR		640	AAC	AAT	N to N							Ü	0.99	£.	0.01

<u> 16. 15</u>

Polymetphism Poly					<u> </u>	SILENT	INT PHI SMS	loa	MISSENSE POLYMORPHISMS	ISE HI SMS	&	NONSENSE POLYMORPHISMS	SE		ALLELE FREQUENCIES	LE NCIE	s
15 642 AC AC T to T T to T T T C 0.96 AC O.96 AC AC T to T T T T T C 0.96 AC O.96 AC AC </td <td>Polymorphism ID</td> <td>вшв</td> <td></td> <td>Codon No.</td> <td>Ref codon</td> <td>Var codon</td> <td></td> <td>Ref codon</td> <td>Var codon</td> <td></td> <td>Ref codon</td> <td>Var codon</td> <td></td> <td>Ref allele</td> <td>Freq</td> <td>Var allele</td> <td>Freq</td>	Polymorphism ID	в шв		Codon No.	Ref codon	Var codon		Ref codon	Var codon		Ref codon	Var codon		Ref allele	Freq	Var allele	Freq
1. DURA CDLR CTT CT	P5u7	F5		642	ACC	ACA	ដ							υ	96.0	4	0.04
12 LDLR CLOR CAS R TO R	LDLRu1	LDLR		653	GTC	GTT	ţ							υ	0.31	Ŧ	0.69
Bu4 ITGA2B CCG CCG<	LDLRu12	LDLR		744	cgg	CGA	to							ຶ່	0.85	4	0.15
Bu4 ITGA2B GCG CCG CCG C P to P GCG GCG CCG CCG <th< td=""><td>ITGA2Bu8</td><td>ITGA2B</td><td></td><td>855</td><td>CTT</td><td>CTC</td><td>ţ</td><td></td><td></td><td></td><td></td><td></td><td></td><td>Į.</td><td>0.99</td><td>υ</td><td>0.01</td></th<>	ITGA2Bu8	ITGA2B		855	CTT	CTC	ţ							Į.	0.99	υ	0.01
Bull Treaze F5 Corrected OFF OFF <t< td=""><td>ITGA2Bu4</td><td>ITGA2B</td><td></td><td>972</td><td>ടാ</td><td>CCA</td><td>t t</td><td></td><td></td><td></td><td></td><td></td><td></td><td>g</td><td>0.99</td><td>-</td><td>0.01</td></t<>	ITGA2Bu4	ITGA2B		972	ടാ	CCA	t t							g	0.99	-	0.01
Horizon Horizon <t< td=""><td>ITGA2Bul</td><td>ITGA2B</td><td></td><td>1021</td><td>GTC</td><td>GTT</td><td>ដ</td><td></td><td></td><td></td><td></td><td></td><td></td><td>ပ</td><td>\vdash</td><td></td><td>0.34</td></t<>	ITGA2Bul	ITGA2B		1021	GTC	GTT	ដ							ပ	\vdash		0.34
1.2 HAGTA H	F5u8	PS												b	-		0.01
u2 HMGCR S9 CF CF CFG LL GD F	HCF2u1	HCF2												ပ		-	0.04
u2 trops trop crop	HMGCRu2	HMGCR												ß	.97	\vdash	0.03
2 CETP 422	ITGB3u2	ITGB3		59				cre	ຄວວ	ដ				4	87	-	0.13
F5 513 613 AAA AAA R to K </td <td>CETPu2</td> <td>CETP</td> <td></td> <td>422</td> <td></td> <td></td> <td></td> <td>ATC</td> <td>GTC</td> <td>ţ٥</td> <td></td> <td></td> <td></td> <td>K</td> <td>34</td> <td>Η</td> <td>99.0</td>	CETPu2	CETP		422				ATC	GTC	ţ٥				K	34	Η	99.0
F5 AT3 CAG CAA CGA CAA	P5u2	PS		513				AGA	AAA	r C				D	-	_	0.15
AT3 CAG CAA Q to Q Q to Q Q 0.62 A Q 0.62 A 5 LDLR 471 AGG AGA R to R Q 0.68 A G 0.68 A LPL LPL 474 A	P5u3	PS		534				CGA	CAA	r C					-	<u> </u>	0.01
5 LDLR 471 AGG AGA R to R G 0.68 A LPL 474 474 C 0.85 C 0.85 C 0.85 G	AT3u2	AT3		337	CAG	CAA	ដ							b			5.38
LPL 474 474 C 0.85 G	LDLRu5	LDLR		471	AGG	AGA	ដ										0.32 `
	LPLu3	Tel		474							TCA	TGA	ţ		-		5.15

FIG. 1E

D-1 TD	
Poly ID	GenBank Acc: Nuc.Position
AT3u1	D29832:1005
AT3u2	D29832:1005
AT3u3	M21645:100
AT3u4	D29832:1374
CETPu1	M30185:1298
CETPu2	M30185:1394
CETPu3	M30185:991
CETPu4	M30185:196
CETPu5	M30185:388
CETPu6	M30185:766
CETPu7	M30185:766
CETPu8	J02898:298
CETPu9	J02898:571
CLanalogul	Z22555:1119
CLanalogu2	Z22555:570
CLanalogu3	Z22555:400
CLanalogu4	Z22555:472
CLanalogu5	Z22555:472 Z22555:972
F2Ru1	M62424:496
F2Ru2	M62424:610
F2Ru3	M62424:664
F2Ru4	M62424:720
F2Ru5	M62424:740
F2Ru6	M62424:405
F2Ru7	M62424:609
F2u1	M17262:10777
F2u2	M17262:15342
F2u3	M17262:15419
F2u4	M17262:13434
F2u5	M17262:16827
F3u1	J02846:9363
F5u1	M14335:1456
F5u2	M14335:1614
F5u3	M14335:1617
F5u4	M14335:1314
F5u5	M14335:1318
F5u6	M14335:1792
F5u7	M14335:2002
HCF2u1	M58600:11907
HCF2u2	M12849:259
HCF2u3	M12849:1353
HCF2u4	M12849:47
HCF2u5	M12849:313
HCF2u6	M12849:651
HCF2u7	M12849:1474
HMGCRu1	M11058:1962
HMGCRu2	M11058:2725

ITGA2Bu2 J02764:2623 ITGA2Bu3 M33320:6845 ITGA2Bu4 J02764:2918 ITGA2Bu5 J02764:2904 ITGA2Bu6 J02764:120 ITGA2Bu7 J02764:2299 ITGA2Bu8 J02764:2567 ITGB3u1 J02703:526 ITGB3u2 J02703:196 ITGB3u3 J02703:902 ITGB3u4 J02703:1163 ITGB3u5 M20311:1549 ITGB3u6 M20311:1561 ITGB3u7 J02703:362 ITGB3u8 J02703:1377 LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1167 LDLRu1 L00347:129 LDLRu1 L00347:129 LDLRu1 L00347:90 LDLRu1 L00347:90 LDLRu1 L00349:107 LDLRu1 L00349:107 LDLRu1 L00345:46 LDLRu4 L00349:44 LDLRu5 L00349:44 LDLRu5 L00349:44 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:534 PTAFRu1 D10202:794 PTAFRu3 D10202:794 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915 TFPIu1 J03225:1006	ITGA2Bu1	M22569:194
ITGA2Bu4	ITGA2Bu2	J02764:2623
ITGA2Bu5	ITGA2Bu3	M33320:6845
ITGA2Bu6	ITGA2Bu4	J02764:2918
ITGA2Bu6	ITGA2Bu5	J02764:2904
ITGA2Bu7 J02764:2299 ITGA2Bu8 J02764:2567 ITGB3u1 J02703:526 ITGB3u2 J02703:196 ITGB3u3 J02703:902 ITGB3u4 J02703:1163 ITGB3u5 M20311:1549 ITGB3u6 M20311:1561 ITGB3u7 J02703:362 ITGB3u8 J02703:1377 LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1167 LDLRu1 L00347:129 LDLRu1 L00347:90 LDLRu1 L00347:90 LDLRu1 L00349:107 LDLRu1 L00349:107 LDLRu1 L00349:44 LDLRu2 L00338:91 LDLRu4 L00349:44 LDLRu5 L00349:44 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1338 LPLu2 M15856:1338 LPLu2 K02059:577 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	ITGA2Bu6	
ITGA2Bu8	ITGA2Bu7	
ITGB3u1	ITGA2Bu8	
ITGB3u2	ITGB3u1	
ITGB3u3	ITGB3u2	
ITGB3u4	ITGB3u3	
ITGB3u5 M20311:1549 ITGB3u6 M20311:1561 ITGB3u7 J02703:362 ITGB3u8 J02703:1377 LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu2 L00349:44 LDLRu3 L00349:44 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	ITGB3u4	J02703:1163
ITGB3u6 M20311:1561 ITGB3u7 J02703:362 ITGB3u8 J02703:1377 LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	ITGB3u5	M20311:1549
ITGB3u8 J02703:1377 LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:783 PTAFRu2 D10202:194 PTAFRu3 D10202:194 PTAFRu4 D10202:195 T	ITGB3u6	
TTGB3u8		
LCATu1 M12625:864 LCATu2 M12625:961 LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00349:107 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00349:44 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M1228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915		
LCATu2 M12625:961 LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00349:46 LDLRu3 L00345:46 LDLRu4 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915		
LCATu3 M12625:1444 LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00349:46 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:577 PROCu2 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LCATu2	
LCATu4 M12625:1167 LDLRu1 L00347:129 LDLRu10 U59436:45 LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00349:46 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M5856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LCATu3	
LDLRu1	LCATu4	
LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00349:44 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu1	
LDLRu11 L00347:90 LDLRu12 L00349:107 LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00349:44 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu10	U59436:45
LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu11	
LDLRu13 L00336:29 LDLRu14 L00351:67 LDLRu2 L00338:91 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu12	L00349:107
LDLRu2 L00338:91 LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu13	
LDLRu3 L00345:46 LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu14	L00351:67
LDLRu4 L00349:44 LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu2	L00338:91
LDLRu5 L00344:70 LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu3	L00345:46
LDLRu6 U59436:93 LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu4	L00349:44
LDLRu7 L29401:691 LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:783 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu5	L00344:70
LDLRu8 L00344:59 LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:783 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu6	U59436:93
LDLRu9 L00343:152 LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:794 PTAFRu3 D10202:783 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu7	L29401:691
LPLu1 M15856:1338 LPLu2 M15856:1453 LPLu3 M76722:3150 PROCu1 K02059:577 PROCu2 K02059:109 PROCu3 M11228:9358 PROCu4 K02059:534 PTAFRu1 D10202:794 PTAFRu2 D10202:1047 PTAFRu3 D10202:783 PTAFRu4 D10202:194 PTAFRu5 D10202:1125 TBXA2Ru1 D38081:1915	LDLRu8	L00344:59
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LOCUS HUMPAFRE 1780 bp mRNA PRI 10-OCT-1992 DEFINITION Human mRNA for platelet-activating factor receptor, complete cds. ACCESSION D10202 D90433 NID g219975 KEYWORDS G-protein coupled receptor; PAF receptor; platelet-activating factor receptor. SOURCE Human leukocytes cDNA to mRNA. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE (bases 1 to 1780) Nakamura, M., Honda, Z., Izumi, T., Sakanaka, C., Mutoh, H., MInami, M., **AUTHORS** Bito, H., Seyama, Y., Noma, M., Mtsumoto, T. and Shimizu, T. TITLE Molecular cloning and expression of platelet-activating factor receptor from human leukocytes JOURNAL J. Biol. Chem. 266 (30), 20400-20405 (1991) MEDLINE 92041873 REFERENCE (bases 1 to 1780) **AUTHORS** Shimizu, T. TITLE Direct Submission **JOURNAL** Submitted (28-JUN-1991) to the DDBJ/EMBL/GenBank databases. Takao Shimizu, Faculty of Medicine, University of Tokyo, Department of Biochemistry; 7-3-1 Hongo, Bunkyo-ku, Tokyo 113, Japan (Tel:03-3812-2111(ex.3448), Fax:03-3813-8732) COMMENT Submitted (28-Jun-1991) to DDBJ by: Takao Shimizu Department of Biochemistry Faculty of Medicine, University of Tokyo 7-3-1 Hongo, Bunkyo-ku Tokyo 113 Japan Phone: 03-3812-2111 x3448 Fax: 03-3813-8732. **FEATURES** Location/Qualifiers source 1..1780 /organism="Homo sapiens" /db_xref="taxon:9606" /cell_type="leukocytes" CDS 113..1141/codon_start=1 /product="platelet-activating factor receptor" /db_xref="PID:d1001519" /db_xref="PID:g219976"

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LOCUS HUMATIIIV 1467 bp mRNA PRI 03-SEP-1996 DEFINITION Human mRNA for antithrombin III variant, complete cds. D29832 ACCESSION g576553 NID KEYWORDS AT-III; antithrombin III. SOURCE Homo sapiens (individual-isolate AT-III Kyoto) cDNA to mRNA, clone pKF16c. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE 1 (sites) **AUTHORS** Tsuji, H., Takada, O., Nakagawa, M., Tanaka, S. and Hashimoto-Gotoh, T. TITLE Hereditary antithrombin III deficiency: identification of an arginine-406 to methionine point mutation near protease reactive site JOURNAL (in) Yoshida, T.O. and Wilson, J.M. (Eds.); MOLECULAR APPROACHES TO THE STUDY AND TREATMENT OF HUMAN DISEASES: 51-55; Elsevier Science (1992) REFERENCE (bases 1 to 1467) AUTHORS Hashimoto-Gotoh, T. JOURNAL Unpublished (1994) FEATURES Location/Qualifiers source 1..1467 /organism="Homo sapiens" /db_xref="taxon:9606" CDS 22..1419 /note="Wild type AT-III has 'g' instead of 't' at position 1337 nt. Also amino acid residue changes from Met to Arg at position 406 aa in wild type AT-III." /codon_start=1 /product="antithrombin III (AT-III) variant" /db_xref="PID:d1006776" /db_xref="PID:g576554" translation="MYSNVIGTVTSGKRKVYLLSLLLIGFWDCVTCHGSPVDICTAKP/ RDIPMNPMCIYRSPEKKATEDEGSEQKIPEATNNRRVWELSKANSRFATTFYQHLADS

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PCT/US99/06473 WO 99/50454

11/97

LOCUS HUMHTAR 2932 bp mRNA PRT 03-APR-1996 Human mRNA for thromboxane A2 receptor, complete cds. DEFINITION ACCESSION D38081 NID g533325 KEYWORDS thromboxane A2 receptor. SOURCE Homo spaiens placenta cDNA to mRNA, clone HPL. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE 1 (bases 1 to 2932) **AUTHORS** Hirata, M., Hayashi, Y., Ushikubi, F., Yokota, Y., Kageyama, R., Nakanishi, S. and Narumiya, S. Cloning and expression of cDNA for a human thromboxane A2 receptor TITLE JOURNAL. Nature 349 (6310), 617-620 (1991) MEDLINE 91156030 REFERENCE 2 (sites) **AUTHORS** Nusing, R.M., Hirata, M., Kakizuka, A., Eki, T., Ozawa, K. and Narumiya, S. TITLE Characterization and chromosomal mapping of the human thromboxane A2 receptor gene J. Biol. Chem. 268 (33), 25253-25259 (1993) JOURNAL 94043399 MEDLINE REFERENCE 3 (bases 1 to 2932) **AUTHORS** Hirata, M. TITLE Direct Submission JOURNAL Submitted (26-AUG-1994) to the DDBJ/EMBL/GenBank databases. Masakazu Hirata, Kyoto University Faculty of Medicine, Department of Pharmacology; Yoshida, Sakyo-ku, Kyoto, Kyoto 606, Japan (Tel:81-75-753-4392, Fax:81-75-753-4693) **FEATURES** Location/Qualifiers source 1..2932 /organism="Homo sapiens" /db_xref="taxon:9606" /tissue_type="placenta" misc_feature 1..705 /note="This part of the cDNA clone may not belong to the thromboxane A2 receptor gene. Please refer to Nuesing, R.M. et al.(reference2)" CDS 992..2023 /codon_start=1 /evidence=experimental /product="Human thromboxane A2 receptor" /db_xref="PID:d1007852" /db_xref="PID:g533326" /translation="MWPNGSSLGPCFRPTNITLEERRLIASPWFAASFCVVGLASNLL

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REFERENCE
              (bases 1 to 3170)
 AUTHORS
            Fitzgerald, L.A., Steiner, B., Rall, S.C. Jr., Lo, S.S. and
            Phillips, D.R.
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COMMENT
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14/97

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LOCUS HUMPLG2B 3303 bp mRNA PRI 07-JAN-1995 DEFINITION Human platelet membrane glycoprotein IIb (ITGA2B) mRNA, complete cds. ACCESSION J02764 NID g190067 KEYWORDS membrane adhesive protein; platelet membrane glycoprotein; platelet receptor. SOURCE Human HEL cell, cDNA to mRNA. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. 1 (bases 1 to 3303) REFERENCE AUTHORS Poncz, M., Eisman, R., Heidenreich, R., Silver, S.M., Vilaire, G., Surrey, S., Schwartz, E. and Bennett, J.S. ጥተጥኒድ Structure of the platelet membrane glycoprotein IIb. Homology to the alpha subunits of the vitronectin and fibronectin membrane receptors JOURNAL J. Biol. Chem. 262 (18), 8476-8482 (1987) MEDLINE 87250457 COMMENT Draft entry and computer-readable sequence [1] kindly provided by M. Poncz, 15-APR-1987. **FEATURES** Location/Qualifiers 1..3303 source /organism="Homo sapiens" /db_xref="taxon:9606" /map="17q21.32" mRNA <1..3303 /gene="ITGA2B" /note="G00-120-012" gene 1..3303 /gene="ITGA2B" sig_peptide 2..94 /gene="ITGA2B" /note="G00-120-012" CDS 2..3121 /gene="ITGA2B" /codon start=1 /db_xref="GDB:G00-120-012" /product="platelet membrane glycoprotein IIb" /db_xref="PID:g190068" /translation="MARALCPLQALWLLEWVLLLLGPCAAPPAWALNLDPVQLTFYAG PNGSQFGFSLDFHKDSHGRVAIVVGAPRTLGPSQEETGGVFLCPWRAEGGQCPSLLFD LRDETRNVGSQTLQTFKARQGLGASVVSWSDVIVACAPWQHWNVLEKTEEAEKTPVGS CFLAQPESGRRAEYSPCRGNTLSRIYVENDFSWDKRYCEAGFSSVVTQAGELVLGAPG GYYFLGLLAQAPVADIFSSYRPGILLWHVSSQSLSFDSSNPEYFDGYWGYSVAVGEFD GDLNTTEYVVGAPTWSWTLGAVEILDSYYQRLHRLRAEQMASYFGHSVAVTDVNGDGR HDLLVGAPLYMESRADRKLAEVGRVYLFLQPRGPHALGAPSLLLTGTOLYGRFGSAIA

FIG. 7A

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            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
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 AUTHORS
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            Complete sequence of the human tissue factor gene, a highly
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  JOURNAL
            Biochemistry 28 (4), 1755-1762 (1989)
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COMMENT
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LOCUS
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                          894 bp
                                    DNA
                                                     PRI
                                                                01-NOV-1994
DEFINITION
            Human cholesteryl ester transfer protein (CETP) gene, exons 15 and
ACCESSION
            M32998 J02898
            g180267
NID
KEYWORDS
            cholesteryl ester transfer protein.
SEGMENT
            7 of 7
SOURCE
            Human DNA
  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 1 to 894)
  AUTHORS
            Agellon, L.B., Quinet, E.M., Gillette, T.G., Drayna, D.T., Brown, M.L.
            and Tall, A.R.
  JOURNAL
            Unpublished (1990)
REFERENCE
            2 (sites)
  AUTHORS
            Agellon, L.B., Quinet, E.M., Gillette, T.G., Drayna, D.T., Brown, M.L.
            and Tall, A.R.
            Organization of the human cholesteryl ester transfer protein gene
  TITLE
  JOURNAL
            Biochemistry 29 (6), 1372-1376 (1990)
  MEDLINE
            90241928
COMMENT
            [2] sites for [1]; intron/exon boundaries.
            Draft entry and computer-readable sequence for [2] kindly
submitted
            by L.B.Agellon, 16-MAR-1990.
FEATURES
                     Location/Qualifiers
     source
                     1..894
                     /organism="Homo sapiens"
                     /db_xref="taxon:9606"
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                     join(M32992:388..1656,M32993:1..3446,M32994:1..628,
                     M32995:1..399,M32996:1..409,M32997:1..1420,1..342)
                     /gene="CETP"
     CDS
                     join(M32992:388..505,M32992:1408..1522,M32993:432..566,
                     M32993:654..724,M32993:954..1041,M32993:2068..2137
                     M32993:2355..2415,M32993:3023..3114,M32994:166..345,
                     M32995:238..288,M32996:128..292,M32997:375..442,
                     M32997:770..803,M32997:1285..1357,257..342,523..597)
                     /note="cholesteryl ester transferase protein precursor"
                     /codon_start=1
                     /db_xref="PID:q180269"
translation="MLAATVLTLALLGNAHACSKGTSHEAGIVCRITKPALLVLNHET/
AKVIQTAFQRASYPDITGEKAMMLLGQVKYGLHNIQISHLSIASSQVELVEAKSIDVS
IQNVSVVFKGTLKYGYTTAWWLGIDQSIDFEIDSAIDLQINTQLTCDSGRVRTDAPDC
YLSFHKLLLHLQGEREPGWIKQLFTNFISFTLKLVLKGQICKEINVISNIMADFVOTR
AASILSDGDIGVDISLTGDPVITASYLESHHKGHFIYKNVSEDLPLPTFSPTLLGDSR
MLYFWFSERVFHSLAKVAFQDGRLMLSLMGDEFKAVLETWGFNTNQEIFQEVVGGFPS
QAQVTVHCLKMPKISCQNKGVVVNSSVMVKFLFPRPDQQHSVAYTFEEDIVTTVQASY
{\tt SKKKLFLSLLDFQITPKTVSNLTESSSESVQSFLQSMITAVGIPEVMSRLEVVFTALM}
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NSKGVSLFDIINPEIITRDGFLLLQMDFGFPEHLLVDFLQSLS"

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prim_transcript <1..772</pre>
                           /note="CETP mRNA and introns"
      intron
                           <1..256
                           /gene="CETP"
                           /note="CETP intron N"
      mat_peptide
                           257..342
                           /gene="CETP"
                           /note="cholesteryl ester transferase protein"
      exon
                           257..342
                           /gene="CETP"
                           /note="G00-119-773"
                           /number=15
      intron
                           343..522
                           /note="CETP intron O"
      exon
                           523..>597
                           /note="cholesteryl ester transferase protein precursor"
                           /number=16
      mat_peptide
                           523..594
                           /note="cholesteryl ester transferase protein"
                           756..762
      polyA_signal
BASE COUNT
                   178 a
                                262 c
                                            256 g
                                                        198 t
               About 950 bp after segment 6.
ORIGIN
        1 ggatgggttg ggagctcaag ttttggggca gaagggaatt ttttttggca gcagagtgca
61 agccctgccg ccaggcaaac tctgctcttc ctcatcctca gaagcacttg ctcactctgc
       121 taaarcaaag tgaaacgcat gtttacagaa tattggtcca aaagggtctc agcatctccc
       181 actacccagg gtgcagagcc tcgggccggc cttgctcccc aagaagggct gactggggct 241 ctgtccctc gcccagggct cgaggtagtg tttacagccc tcatgaacag caaaggcgtg
       301 agectetteg acateateaa ecetgagatt ateaetegag atgtgagtae aaageceeee
       361 tcaccagccc ctgttcctgg ggagagaggc ccagacagga ttcctggggt gactgggggc 421 tgttggggag acagacagag gggcctctac cagcttggct ccctcctggt ggcctgggag
       481 tragercage tegeretet etectactge ecetecette agggetteet getgetgeag
       541 atggactttg getteeetga geacetgetg gtggatttee tecagagett gagetagaag
601 tetecaagga ggtegggatg gggettgtag cagaaggeaa geaceagget cacagetgga
       661 accetggtgt cteeteeage gtggtggaag ttgggttagg agtacggaga tggagattgg
       721 ctcccaactc ctccctatcc taaaggccca ctggcattaa agtgctgtat ccaagagctg
781 cggagtcctt cttctgtggc tggcgggtag agggggggg aagggattgt ctcaccagtg
       841 ccgtccacct cttttcagcc cttccaagca gctgccccca aaccctccaa gctt
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LOCUS HUMCILA 1431 bp mRNA PRI 01-NOV-1994 DEFINITION Human lipoprotein-associated coagulation inhibitor mRNA, complete cds. ACCESSION J03225 NID g180545 KEYWORDS lipoprotein-associated coagulation inhibitor. SOURCE Human placenta, cDNA to mRNA, clone lambda-P9. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. 1 (bases 1 to 1431) REFERENCE **AUTHORS** Wun, T.C., Kretzmer, K.K., Girard, T.J., Miletich, J.P. and Broze, G.J. Jr. TITLE Cloning and characterization of a cDNA coding for the lipoprotein-associated coagulation inhibitor shows that it consists of three tandem Kunitz-type inhibitory domains JOURNAL. J. Biol. Chem. 263 (13), 6001-6004 (1988) 88198127 MEDLINE COMMENT Draft entry and printed copy of sequence for [1] kindly provided ÒΥ T.-C.Wun, 19-MAR-1988. **FEATURES** Location/Qualifiers source 1..1431 /organism="Homo sapiens" /db_xref="taxon:9606" /map="2q31-q32.1" sig_peptide 133..216 /gene="TFPI" /note="lipoprotein-associated coagulation inhibitor signal peptide* CDS 133..1047 /gene="TFPI" /note="lipoprotein-associated coagulation inhibitor precursor /codon_start=1 /db_xref="GDB:G00-127-364" /db_xref="PID:g180546"

/translation="MIYTMKKVHALWASVCLLLNLAPAPLNADSEEDEEHTIITDTEL

 ${\tt PPLKLMHSFCAFKADDGPCKAIMKRFFFNIFTRQCEEFIYGGCEGNQNRFESLEECKK}$

MCTRDNANRIIKTTLQQEKPDFCFLEEDPGICRGYITRYFYMNQTKQCERFKYGGCLG

 ${\tt NMNNFETLEECKNICEDGPNGFQVDNYGTQLNAVNNSLTPQSTKVPSLFEFHGPSWCL}$

TPADRGLCRANENRFYYNSVIGKCRPFKYSGCGGNENNFTSKQECLRACKKGFIQRIS KGGLIKTKRKKKQRVKIAYEEIFVKNM*

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gene
                          133..1047
                          /gene="TFPI"
      mat_peptide
                          217..1044
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                          /note="lipoprotein-associated coagulation inhibitor"
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BASE COUNT
                              244 c
                                        267 g
                                                     441 t
ORIGIN
              351 bp upstream of SspI site.
         1 ggcgggtctg cttctaaaag aagaagtaga gaagataaat cctgtcttca atacctggaa
        61 ggaaaaacaa aataacctca actccgtttt gaaaaaaaca ttccaagaac tttcatcaga
       121 gattttactt agatgattta cacaatgaag aaagtacatg cactttgggc ttctgtatgc 181 ctgctgctta atcttgcccc tgcccctctt aatgctgatt ctgaggaaga tgaagaacac
       241 acaattatca cagatacgga gttgccacca ctgaaactta tgcattcatt ttgtgcattc
       301 aaggeggatg atggeecatg taaageaate atgaaaagat ttttetteaa tatttteaet 361 egacagtgeg aagaatttat atatggggga tgtgaaggaa ateagaateg atttgaaagt
       421 ctggaagagt gcaaaaaaat gtgtacaaga gataatgcaa acaggattat aaagacaaca
       481 ttgcaacaag aaaagccaga tttctgcttt ttggaagaag atcctggaat atgtcgaggt
       541 tatattacca ggtattttta taacaatcag acaaaacagt gtgaacgttt caagtatggt
       601 ggatgcctgg gcaatatgaa caattttgag acactggaag aatgcaagaa catttgtgaa
       661 gatggtccga atggtttcca ggtggataat tatggaaccc agctcaatgc tgtgaataac
721 tccctgactc cgcaatcaac caaggttccc agcctttttg aatttcacgg tccctcatgg
       781 tgtctcactc cagcagacag aggattgtgt cgtgccaatg agaacagatt ctactacaat
       841 tcagtcattg ggaaatgccg cccatttaag tacagtggat gtgggggaaa tgaaaacaat
901 tttacttcca aacaagaatg tctgagggca tgtaaaaaaag gtttcatcca aagaatatca
961 aaaggaggcc taattaaaac caaaagaaaa agaaagaagc agagagtgaa aatagcatat
      1021 gaagaaattt ttgttaaaaa tatgtgaatt tgttatagca atgtaacatt aattctacta
      1081 aatattttat atgaaatgtt tcactatgat tttctatttt tcttctaaaa tcgttttaat
      1141 taatatgttc attaaatttt ctatgcttat tgtacttgtt atcaacacgt ttgtatcaga
      1201 gttgcttttc taatcttgtt aaattgctta tictaggict gtaatttatt aactggctac
      1261 tgggaaatta cttattttct ggatctatct gtattttcat ttaactacaa attatcatac
      1321 taccggctac atcaaatcag teetttgatt ceatttggtg accatetgtt tgagaatatg
      1381 atcatgtaaa tgattatctc ctttatagcc tgtaaccaga ttaagccccc c
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27/97

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LOCUS
           HUMPRC
                         1366 bp
                                    mRNA
                                                     PRI
                                                               08-JAN-1995
DEFINITION
           Human protein C, mRNA.
ACCESSION
            K02059
            g190322
NID
KEYWORDS
            glycoprotein; protease; protein C; serine protease.
SOURCE
            Human liver, cDNA (library of Woo) to mRNA, clones lambda-HC1026
            and lambda-HC1375.
  ORGANISM
           Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 1 to 1366)
  AUTHORS
            Foster, D. and Davie, E.W.
            Characterization of a cDNA coding for human protein C
  TITLE
  JOURNAL
            Proc. Natl. Acad. Sci. U.S.A. 81 (15), 4766-4770 (1984)
  MEDLINE
            84272714
            Protein C is a precursor to a serine protease called 'activated
COMMENT
            protein C' that has a strong anticoagulant activity. The amino
acid
            sequence as determined from the cDNA indicates that protein {\tt C} is
            synthesized as a single-chain polypeptide containing the light
            chain and the heavy chain connected by a dipeptide of Lys-Arg.
This
            precursor peptide is then converted to the light and heavy chains
            by cleavage of two or more internal peptide bonds. The amino acid
            sequence of human protein C shows a high homology with that of the
            bovine molecule. Two clones were sequenced in [1] and shown to
            code for human protein C. Clone lambda-HC1026 covers bp 146-1140,
            and clone lambda-HC1375 covers bp 1-1366.
                                                        The two cDNA clones had
            a poly-A tail at different positions; both poly-A sites were
            preceded by poly-A signals [1].
FEATURES
                     Location/Qualifiers
    source
                     1..1366
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                     /db_xref="taxon:9606"
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                     /tissue_lib="of Woo"
                     /map="2q13-q21"
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                     <1..1366
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                     /note="G00-120-317"
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                     <1..1140
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                     /note="G00-120-317"
     gene
                     1..1366
                     /gene="PROC"
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                      /note="G00-120-317"
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                      /gene="PROC"
                     /note="."
                      /codon_start=2
                      /db_xref="GDB:G00-120-317"
                      /product="protein C"
                     /db_xref="PID:g190323"
/translation="QGHGTCIDGIGSFSCDCRSGWEGRFCQREVSFLNCSLDNGGCTH
YCLEEVGWRRCSCAPGYKLGDDLLQCHPAVKFPCGRPWKRMEKKRSHLKRDTEDOEDO
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FIG. 11A

VDPRLIDGKMTRRGDSPWQVVLLDSKKKLACGAVLIHPSWVLTAAHCMDESKKLLVRL GEYDLRRWEKWELDLDIKEVFVHPNYSKSTTDNDIALLHLAQPATLSQTIVPICLPDS

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GLAERELNQAGQETLVTGWGYHSSREKEAKRNRTFVLNFIKIPVVPHNECSEVMSNMV
SENMLCAGILGDRQDACEGDSGGPMVASFHGTWFLVGLVSWGEGCGLLHNYGVYTKVS
                          RYLDWIHGHIRDKEAPOKSWAP*
      mat_peptide
                          284..1069
                          /gene="PROC"
                           /note="G00-120-317"
                          /product="protein C heavy chain"
      mat_peptide
                          320..1069
                          /gene="PROC"
                          /note="G00-120-317"
                          /product="protein C activated heavy chain"
BASE COUNT
                    302 a
                                          425 g
                               388 c
                                                      251 t
               207 bp upstream of PstI site; chromosome 2q14-q21.
ORIGIN
          1 ccaagggcac ggcacgtgca tcgacggcat cggcagcttc agctgcgact gccgcagcgg
       61 ctgggagggc cgcttctgcc agcgcgaggt gagcttcctc aattgctctc tggacaacgg
121 cggctgcacg cattactgcc tagaggaggt gggctggcgg cgctgtagct gtgcgcctgg
       181 ctacaagetg ggggacgace teetgeagtg teaceeegea gtgaagttee ettgtgggag
       241 gccctggaag cggatggaga agaagcgcag tcacctgaaa cgagacacag aagaccaaga
301 agaccaagta gatccgcggc tcattgatgg gaagatgacc aggcggggag acagcccctg
       361 gcaggtggtc ctgctggact caaagaagaa gctggcctgc ggggcagtgc tcatccaccc
       421 ctcctgggtg ctgacagcgg cccactgcat ggacgagtcc aagaagctcc ttgtcaggct
       481 tggagagtat gacctgcggc gctgggagaa gtgggagctg gacctggaca tcaaggaggt
541 cttcgtccac cccaactaca gcaagagcac caccgacaat gacatcgcac tgctgcacct
       601 ggcccagccc gccaccctct cgcagaccat agtgcccatc tgcctcccgg acagcggcct
       661 tgcagagcgc gagctcaatc aggccggcca ggagaccctc gtgacgggct ggggctacca
721 cagcagccga gagaaggagg ccaagagaaa ccgcaccttc gtcctcaact tcatcaagat
       781 tecegtggte eegcacaatg agtgeagega ggteatgage aacatggtgt etgagaacat
       841 gctgtgtgcg ggcatcctcg gggaccggca ggatgcctgc gagggcgaca gtggggggcc
901 catggtcgcc tccttccacg gcacctggtt cctggtgggc ctggtgagct ggggtgaggg
       961 ctgtgggctc cttcacaact acggcgttta caccaaagtc agccgctacc tcgactggat
      1021 ccatgggcac atcagagaca aggaagcccc ccagaagagc tgggcacctt agcgaccctc 1081 cctgcagggc tgggcttttg catggcaatg gatgggacat taaagggaca tgtaacaagc
      1141 acaccggcct gctgttctgt ccttccatcc ctcttttggg ctcttctgga gggaagtaac
      1201 atttactgag cacctgttgt atgtcacatg ccttatgaat agaatcttaa ctcctagagc
      1261 aactctgtcg ggtggggagg agcagatcca agttttgcgg ggtctaaagc tgtgtgtgtt
      1321 gagggggata ctctgtttat gaaaaagaat aaaaaacaca accacg
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```
LOCUS
            HUMLDLR02
                           144 bp
                                     DNA
                                                      PRI
                                                                30-NOV-1994
DEFINITION Human low density lipoprotein receptor gene, exon 2.
            L00336 K02573
ACCESSION
            g187078
NID
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
 ORGANISM
           Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 16 to 138)
 AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
 TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
            Cell 39 (1), 27-38 (1984)
 JOURNAL
            85024898
 MEDLINE
REFERENCE
               (bases 1 to 23; 132 to 144)
 AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
 TITLE
            The LDL receptor gene: a mosaic of exons shared with different
 JOURNAL
            Science 228 (4701), 815-822 (1985)
 MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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     exon
                     16..138
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                      /number=2
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                     139..>144
                      /gene="LDLR"
                      /note="LDL intron B"
BASE COUNT
            33 a 33 c 46 g 32 t Chromosome 19p13.2-p13.1; about 10 kb after segment 1.
ORIGIN
        1 tttcctctct ctcagtgggc gacagatgtg aaagaaacga gttccagtgc caagacggga
       61 aatgcatctc ctacaagtgg gtctgcgatg gcagcgctga gtgccaggat ggctctgatg
      121 agtcccagga gacgtgctgt gagt
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LOCUS
            HUMLDLR04
                           402 bp
                                      DNA
                                                        PRI
                                                                   30-NOV-1994
DEFINITION Human low density lipoprotein receptor gene, exon 4.
ACCESSION
            L00338 K02573
NID
            g187080
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
            4 of 18
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
            1 (bases 16 to 396)
REFERENCE
  AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
  JOURNAL
            Cell 39 (1), 27-38 (1984)
            85024898
  MEDLINE
REFERENCE
               (bases 1 to 23; 389 to 402)
  AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  TITLE
            The LDL receptor gene: a mosaic of exons shared with different
            proteins
  JOURNAL
            Science 228 (4701), 815-822 (1985)
  MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                      /map="19p13.3"
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                      /gene="LDLR"
                      /note="LDL intron C"
     exon
                      16.:396
                      /gene="LDLR"
                      /note="G00-119-362"
                      /number=4
     intron
                      397..>402
                      /gene="LDLR"
                      /note="LDL intron D"
BASE COUNT
            73 a 131 c 120 g 78 t Chromosome 19p13.2-p13.1; about 2.4 kb after segment 3.
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      61 agtgcatete teggeagtte gtetgtgaet cagaceggga etgettggae ggeteagaeg
121 aggeeteetg eeeggtgete acetgtggte eegeeagett eeagtgeaac ageteeacet
      181 gcatccccca gctgtgggcc tgcgacaacg accccgactg cgaagatggc tcggatgagt
      241 ggccgcagcg ctgtaggggt ctttacgtgt tccaagggga cagtagcccc tgctcggcct
      301 tcgagttcca ctgcctaagt ggcgagtgca tccactccag ctggcgctgt gatggtggcc
      361 ccgactgcaa ggacaaatct gacgaggaaa actgcggtat gg
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```
LOCUS
            HUMLDLR09
                          . 193 bp
                                      DNA
                                                                   30-NOV-1994
                                                        PRI
DEFINITION Human low density lipoprotein receptor gene, exon 9.
            L00343 K02573
ACCESSION
            g187085
NID
            low density lipoprotein receptor-1; repeat region.
KEYWORDS
SEGMENT
            9 of 18
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
             [1].
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 16 to 187)
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
  AUTHORS
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
           sequences in its mRNA
            Cell 39 (1), 27-38 (1984)
  JOURNAL
  MEDLINE
            85024898
            2 (bases 1 to 23; 180 to 193)
REFERENCE
  AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  TITLE
            The LDL receptor gene: a mosaic of exons shared with different
            proteins
  JOURNAL.
            Science 228 (4701), 815-822 (1985)
  MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                      /db_xref="taxon:9606"
                      /map="19p13.3"
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/note="LDL intron H"
                      16..187
     exon
                      /gene="LDLR"
                      /note="G00-119-362"
                      /number=9
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BASE COUNT
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                          64 c
                                     52 g
                                               33 t
ORIGIN
            Chromosome 19p13.2-p13.1; about 1.2 kb after segment 8.
        1 tecceggace eccaggetee ategeetace tettetteac caaceggeac gaggteagga
      61 agatgacget ggaceggage gagtacacca geeteateee caacetgagg aacgtggteg 121 etetggacae ggaggtggee ageaatagaa tetaetggte tgaeetgtee cagagaatga
      181 tctgcaggtg agc
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```
LOCUS
             HUMLDLR10
                             249 bp
                                         DNA
                                                            PRI
                                                                       30-NOV-1994
DEFINITION Human low density lipoprotein receptor gene, exon 10.
ACCESSION
             L00344 K02573
NID
             g187086
KEYWORDS
             low density lipoprotein receptor-1; repeat region.
SEGMENT
             10 of 18
SOURCE
             Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
              [1].
  ORGANISM
             Homo sapiens
             Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
             Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
             1 (bases 16 to 243)
  AUTHORS
             Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
             Goldstein, J.L. and Russell, D.W.
  TITLE
             The human LDL receptor: a cysteine-rich protein with multiple Alu
             sequences in its mRNA
             Cell 39 (1), 27-38 (1984)
  JOURNAL
  MEDLINE
             85024898
REFERENCE
                 (bases 1 to 23; 236 to 249)
  AUTHORS
             Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  TITLE
             The LDL receptor gene: a mosaic of exons shared with different
             proteins
  JOURNAL
             Science 228 (4701), 815-822 (1985)
  MEDLINE
             85218750
COMMENT
             Draft entry and computer-readable sequence for [1] kindly provided
             by D.Russell, 01-MAR-1985.
FEATURES
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                        /map=*19p13.3*
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                        /note="LDL intron I"
     exon
                        16..243
                        /gene="LDLR"
                        /note="G00-119-362"
                        /number=10
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                        244..>249
                        /gene="LDLR"
                        /note="LDL intron J"
BASE COUNT
             51 a 77 c 71 g 50 t
Chromosome 19p13.2-p13.1; about 900 bp after segment 9.
ORIGIN
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      61 catcagcagg gacatccagg ccccgacgg gctggctgtg gactggatcc acagcaacat 121 ctactggacc gactctgtcc tgggcactgt ctctgttgcg gataccaagg gcgtgaagag 181 gaaaacgtta ttcagggaga acggctcaa gccaagggcc atcgtggtgg atcctgttca 241 tgggtgcgt
```

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LOCUS
            HUMLDLR11
                          140 bp
                                     DNA
                                                      PRT
                                                                30-NOV-1994
DEFINITION
            Human low density lipoprotein receptor gene, exon 11.
            L00345 K02573
ACCESSION
NID
            g187087
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
            11 of 18
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 6 to 134)
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
  AUTHORS
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
  JOURNAL.
            Cell 39 (1), 27-38 (1984)
  MEDLINE
            85024898
REFERENCE
            2 (bases 1 to 22; 128 to 140)
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  AUTHORS
            The LDL receptor gene: a mosaic of exons shared with different
  TITLE
            proteins
  JOITENAL.
            Science 228 (4701), 815-822 (1985)
  MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                     /map="19p13.3"
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                     /note="LDL intron J"
     exon
                     16..134
                     /gene="LDLR"
                     /note="G00-119-362"
                     /number=11
     intron
                     135..>140
                     /gene="LDLR"
                     /note="LDL intron K"
                          38 c
BASE COUNT
                 34 a
                                   37 g
                                             31 t
ORIGIN
           Chromosome 19p13.2-p13.1; about 2.6 kb after segment 10.
       1 ctgtcctccc accagcttca tgtactggac tgactgggga actcccgcca agatcaagaa
       61 aggggggcctg aatggtgtgg acatctactc gctggtgact gaaaacattc agtggcccaa
      121 tggcatcacc ctaggtatgt
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LOCUS
            HUMLDLR13
                           163 bp
                                     DNA
                                                      PRI
                                                                 30-NOV-1994
DEFINITION Human low density lipoprotein receptor gene, exon 13.
ACCESSION
            L00347 K02573
NID
            g187089
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
            13 of 18
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
             [1].
  ORGANISM
           Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
            1 (bases 16 to 157)
REFERENCE
  AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
            Cell 39 (1), 27-38 (1984)
  JOURNAL
            85024898
 MEDLINE
REFERENCE
               (bases 1 to 24; 151 to 163)
  AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  TITLE
            The LDL receptor gene: a mosaic of exons shared with different
            proteins
  JOURNAL
            Science 228 (4701), 815-822 (1985)
 MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                      /map="19p13.3"
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                      <1..15
                      /gene="LDLR"
                      /note="LDL intron L"
     exon
                      16..157
                      /gene="LDLR"
                      /note="G00-119-362"
                      /number=13
     intron
                     158..>163
                      /gene="LDLR"
                      /note="LDL intron M"
BASE COUNT
            43\ a 45\ c 34\ g 41\ t Chromosome 19p13.2-p13.1; about 3 kb after segment 12.
ORIGIN
        1 ttgctgcctg tttaggacaa agtattttgg acagatatca tcaacgaagc cattttcagt
       61 gccaaccgcc tcacaggttc cgatgtcaac ttgttggctg aaaacctact gtccccagag
      121 gatatggtcc tcttccacaa cctcacccag ccaagaggta agg
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LOCUS
            HUMLDLR15
                           192 bp
                                     DNA
                                                                30-NOV-1994
                                                      PRI
DEFINITION Human low density lipoprotein receptor gene, exon 15.
ACCESSION
            L00349 K02573
NID
            g187091
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
            15 of 18
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 16 to 186)
  AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
            Cell 39 (1), 27-38 (1984)
  JOURNAL
 MEDLINE
            85024898
               (bases 1 to 23; 179 to 192)
REFERENCE
  AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
            The LDL receptor gene: a mosaic of exons shared with different
  TITLE
            proteins
  JOURNAL
            Science 228 (4701), 815-822 (1985)
 MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                      /db_xref="taxon:9606"
                      /map="19p13.3"
     intron
                      <1..15
                      /gene="LDLR"
                      /note="LDL intron N"
     exon
                      16..186
                      /gene="LDLR"
                      /note="G00-119-362"
                      /number=15
     intron
                      187..>192
                      /gene="LDLR"
                      /note="LDL intron O"
BASE COUNT
            46 a 64 c 49 g 33 t Chromosome 19p13.2-p13.1; about 2.8 kb after segment 14.
ORIGIN
        1 tatttattct ttcagaggct gaggctgcag tggccaccca ggagacatcc accgtcaggc
       61 taaaggtcag ctccacagcc gtaaggacac agcacacaac cacceggcct gttcccgaca
      121 cctcccggct gcctggggcc acccctgggc tcaccacggt ggagatagtg acaatgtctc
      181 accaaggtaa ag
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```
LOCUS
            HUMLDLR17
                           179 bp
                                     DNA
                                                                 30-NOV-1994
                                                      PRI
DEFINITION Human low density lipoprotein receptor gene, exon 17.
ACCESSION
            L00351 K02573
            g187093
NID
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
SEGMENT
            17 of 18
SOURCE
            Human DNA [3] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
               (bases 16 to 173)
  AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
  JOURNAL.
            Cell 39 (1), 27-38 (1984)
  MEDLINE
            85024898
REFERENCE
               (bases 57 to 101)
  AUTHORS
            Lehrman, M.A., Goldstein, J.L., Brown, M.S., Russell, D.W. and
            Schneider, W.J.
  TITLE
            Internalization-defective LDL receptors produced by genes with
            nonsense and frameshift mutations that truncate the cytoplasmic
            domain
  JOURNAL
            Cell 41 (3), 735-743 (1985)
  MEDLINE
            85228224
REFERENCE
               (bases 1 to 23; 164 to 179)
  AUTHORS
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  TITLE
            The LDL receptor gene: a mosaic of exons shared with different
            proteins
  JOURNAL
            Science 228 (4701), 815-822 (1985)
  MEDLINE
            85218750
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
FEATURES
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                      /note="LDL intron P"
     exon
                      16..173
                      /gene="LDLR".
                      /note="G00-119-362"
                      /number=17
     mutation
                      76..77
                      /gene="LDLR"
                      /note="ac in wt; aagaac in internalization-defective
                      familial hypercholesterolemia [2]"
     intron
                     174..>179
                      /gene="LDLR"
                      /note="LDL intron Q"
            42 a 56 c 39 g 42 t Chromosome 19p13.2-p13.1; about 1.4 kb after segment 16.
BASE COUNT
ORIGIN
        1 tgcctctccc tacagigctc ctcgtcttcc tttgcctggg ggtcttcctt ctatggaaga
       61 actggcggct taagaacatc aacagcatca actttgacaa ccccgtctat cagaagacca
     121 cagaggatga ggtccacatt tgccacaacc aggacggcta cagctacccc tcggtgagt
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WO-99/50454 PCT/US99/06473

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LOCUS
            HUMLDLR01
                           769 bp
                                      DNA
                                                       PRI
                                                                  30-NOV-1994
DEFINITION
            Human low density lipoprotein receptor gene, exon 1.
ACCESSION
            L29401 K02573 M10664 N00033
NID
            q460288
KEYWORDS
            low density lipoprotein receptor-1; repeat region.
            1 of 18
SEGMENT
SOURCE
            Human DNA [2] and fetal adrenal gland, cDNA to mRNA, clone pLDLR-2
            [1].
  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (sites)
  AUTHORS
            Yamamoto, T., Davis, C.G., Brown, M.S., Schneider, W.J., Casey, M.L.,
            Goldstein, J.L. and Russell, D.W.
  TITLE
            The human LDL receptor: a cysteine-rich protein with multiple Alu
            sequences in its mRNA
            Cell 39 (1), 27-38 (1984)
  JOURNAL
            85024898
  MEDLINE
REFERENCE
               (bases 1 to 769)
            Sudhof, T.C., Goldstein, J.L., Brown, M.S. and Russell, D.W.
  AUTHORS
  TITLE
            The LDL receptor gene: a mosaic of exons shared with different
  JOURNAL
            Science 228 (4701), 815-822 (1985)
  MEDITNE
            85218750
            Bases 1-769 from Science 228, 815-822 (1985)
Bases 675-754 from Cell 39, 27-38 (1984)
COMMENT
            Draft entry and computer-readable sequence for [1] kindly provided
            by D.Russell, 01-MAR-1985.
                      Location/Qualifiers
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                      /map="19p13.3"
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                      /note="low density lipoprotein receptor; G00-119-362"
                      /number=1
     sig_peptide
                      688..750
                      /gene="LDLR"
                      /note="low density lipoprotein receptor signal pept"
     intron
                      755..>769
                      /gene="LDLR"
                      /note="LDL intron A"
                          169 c
BASE COUNT
                                    194 g
                                             ·186 t
ORIGIN
            Chromosome 19p13.2-p13.1; 1 bp upstream of BamHI site.
       1 ggatcccaca aaacaaaaaa tatttttttg gctgtacttt tgtgaagatt ttatttaaat 61 tcctgattga tcagtgtcta ttaggtgatt tggaataaca atgtaaaaac aatatacaac
      121 gaaaggaagc taaaaatcta tacacaattc ctagaaagga aaaggcaaat atagaaagtg
      181 gcggaagttc ccaacatttt tagtgttttc cttttgaggc agagaggaca atggcattag
      241 gctattggag gatcttgaaa ggctgttgtt atccttctgt ggacaacaac agcaaaatgt
      301 taacagttaa acatcgagaa atttcaggag gatctttcag aagatgcgtt tccaattttg
      361 agggggcgtc agctettcac eggagaceca aatacaacaa atcaagtege etgeeetgge
      421 gacactttcg aaggactgga gtgggaatca gagcttcacg ggttaaaagc cgatgtcaca
      481 teggeegtte gaaacteete etettgeagt gaggtgaaga catttgaaaa teacceeact
      541 gcaaactcct ccccctgcta gaaacctcac attgaaatgc tgtaaatgac gtgggccccg
      601 agtgcaatcg cgggaagcca gggtttccag ctaggacaca gcaggtcgtg atccgggtcg
      661 ggacactgcc tggcagaggc tgcgagcatg gggccctggg gctggaaatt gcgctggacc
      721 gtcgccttgc tcctcgccgc ggcggggact gcaggtaagg cttgctcca
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LOCUS
            HUMF511
                           279 bp
                                      DNA
                                                       PRI
                                                                  10-NOV-1994
DEFINITION Human coagulation factor V gene, exon 11.
ACCESSION
            L32765 J05368
NID
            q488094
KEYWORDS
            coagulation factor V; factor V.
SEGMENT
            11 of 25
SOURCE
            Homo sapiens DNA.
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 1 to 279)
            Kane, W.H. and Davie, E.W.
  AUTHORS
  TITLE
            Cloning of a cDNA coding for human factor V, a blood coagulation
            factor homologous to factor VIII and ceruloplasmin
  JOURNAL
            Proc. Natl. Acad. Sci. U.S.A. 83 (18), 6800-6804 (1986)
  MEDLINE
            86313665
REFERENCE
            2 (bases 1 to 279)
            Kane, W.H., Ichinose, A., Hagen, F.S. and Davie, E.W.
  AUTHORS
            Cloning of cDNAs coding for the heavy chain region and connecting
  TITLE
            region of human factor V, a blood coagulation factor with four
            types of internal repeats
            Biochemistry 26 (20), 6508-6514 (1987)
  JOURNAL
  MEDLINE
            88107560
REFERENCE
               (bases 1 to 279)
            Jenny, R.J., Pittman, D.D., Toole, J.J., Kriz, R.W., Aldape, R.A.,
  AUTHORS
            Hewick, R.M., Kaufman, R.J. and Mann, K.G.
  TITLE
            Complete cDNA and derived amino acid sequence of human factor V
            Proc. Natl. Acad. Sci. U.S.A. 84 (14), 4846-4850 (1987)
  JOURNAL.
  MEDLINE
            87260886
REFERENCE
               (bases 1 to 279)
            Cripe, L.D., Moore, K.D. and Kane, W.H.
  AUTHORS
  TITLE
            Structure of the gene for human coagulation factor V
  JOURNAL
            Biochemistry 31 (15), 3777-3785 (1992)
  MEDLINE
            92232668
            5 (bases 1 to 279)
REFERENCE
            Shen, N.L., Fan, S.T., Pyati, J., Graff, R., LaPolla, R.J. and
  AUTHORS
            Edgington, T.S.
  TITLE
            The serine protease cofactor factor V is synthesized by
lymphocytes
  JOURNAL
            J. Immunol. 150 (7), 2992-3001 (1993)
            93203619
  MEDLINE
FEATURES
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                      /cell_type="fibroblast"
                      /map="1q21-q25"
                      order(L32764:277..>319,<1..74)
     intron
                      /gene="F5
                      /note="3.1 kb gap; G00-119-896"
                      /number=10
     exon
                      75..225
                      /gene="F5"
                      /note="G00-119-896"
                      /number=11
BASE COUNT
                 73 a
                           52 c
                                     61 g
                                               93 t
ORIGIN
        1 tctgagttct ctattctgtt ccattggtct atgcgtctgt tcttgtacca gtactatact
      61 gttttgtcct ccagagggca gcagacatcg aacagcaggc tgtgtttgct gtgtttgatg
121 agaacaaaag ctggtacctt gaggacaaca tcaacaagtt ttgtgaaaat cctgatgagg
      181 tgaaacgtga tgaccccaag ttttatgaat caaacatcat gagcagtaag tcagagtact
      241 attitigtic atcagtitti catticity gitgaaata
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WO 99/50454 PCT/US99/06473

39/97

LOCUS HUMHMGCOA 2904 bp mRNA PRI 08-NOV-1994 DEFINITION Human 3-hydroxy-3-methylglutaryl coenzyme A reductase mRNA, complete cds. ACCESSION M11058 g184243 NID KEYWORDS 3-hydroxy-3-methylglutaryl coenzyme A reductase; glycoprotein. SOURCE Human fetal adrenal gland, cDNA to mRNA, library of T.Maniatis, clone pHRed-102. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE (bases 1 to 2904) AUTHORS Luskey, K.L. and Stevens, B. Human 3-hydroxy-3-methylglutaryl coenzyme A reductase. Conserved TITLE domains responsible for catalytic activity and sterol-regulated degradation JOURNAL J. Biol. Chem. 260 (18), 10271-10277 (1985) MEDLINE 85261451 COMMENT Draft entry and sequence in computer readable form for [1] kindly provided by K.L.Luskey, 16-JAN-1986. HMG-CoA reductase is the rate-limiting enzyme for cholesterol synthesis and is regulated via a negative feedback mechanism mediated by sterols and non-sterol metabolites derived from mevalonate, the product of the reaction catalyzed by reductase. Normally in mammalian cells this enzyme is suppressed by cholesterol derived from the internalization and degradation of low density lipoprotein (LDL) via the LDL receptor. Competitive inhibitors of the reductase induce the expression of LDL receptors in the liver, which in turn increases the catabolism of plasma LDL and lowers the plasma concentration of cholesterol, an important determinant of atherosclerosis. The sequence coding for the highly conserved membrane bound region of the protein is located at positions 51-1067, that coding for the linker part of the protein at positions 1068-1397 and for the strongly conserved water-soluble catalytic part at positions 1398-2714. **FEATURES** Location/Qualifiers source 1..2904 /organism="Homo sapiens" /db_xref="taxon:9606" /map="5q13.3-q14" mRNA <1..>2904 /note="HMG CoA mRNA" gene 51..2717 /gene="HMGCR" CDS 51..2717 /gene="HMGCR" /note="3-hydroxy-3-methylglutaryl coenzyme A reductase" /codon_start=1 /db_xref="GDB:G00-119-312" /db_xref=*PID:g306865

/translation="MLSRLFRMHGLFVASHPWEVIVGTVTLTICMMSMNMFTGNNKIC

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SFVFSTVVIHFLDKELTGLNEALPFFLLLIDLSRASTLAKFALSSNSQDEVRENIARG

MAILGPTFTLDALVECLVIGVGTMSGVRQLEIMCCFGCMSVLANYFVFMTFFPACVSL

VLELSRESREGRPIWQLSHFARVLEEEENKPNPVTQRVKMIMSLGLVLVHAHSRWIAD

FIG. 22A

```
PSPQNSTADTSKVSLGLDENVSKRIEPSVSLWQFYLSKMISMDIEQVITLSLALLLAV
KYIFFEQTETESTLSLKNPITSPVVTQKKVPDNCCRREPMLVRNNQKCDSVEEETGIN
RERKVEVIKPLVAETDTPNRATFVVGNSSLLDTSSVLVTQEPEIELPREPRPNEECLQ
ILGNAEKGAKFLSDAEIIQLVNAKHIPAYKLETLMETHERGVSIRRQLLSKKLSEPSS
LQYLPYRDYNYSLVMGACCENVIGYMPIPVGVAGPLCLDEKEFQVPMATTEGCLVAST
NRGCRAIGLGGGASSRVLADGMTRGPVVRLPRACDSAEVKAWLETSEGFAVIKEAFDS
TSRFARLQKLHTSIAGRNLYIRFQSRSGDAMGMNMISKGTEKALSKLHEYFPEMQILA
VSGNYCTDKKPAAINWIEGRGKSVVCEAVIPAKVVREVLKTTTEAMIEVNINKNLVGS
AMAGSIGGYNAHAANIVTAIYIACGQDAAQNVGSSNCITLMEASGPTNEDLYISCTMP
SIEIGTVGGGTNLLPQQACLQMLGVQGACKDNPGENARQLARIVCGTVMAGELSLMAA
                      LAAGHLVKSHMIHNRSKINLQDLQGACTKKTA"
BASE COUNT
                          597 c
                                  678 g
                                              807 t
ORIGIN
            27 bp upstream of BamHI site; chromosome 5q13.3-q14.
        1 ttcggtggcc tctagtgaga tctggaggat ccaaggattc tgtagctaca atgttgtcaa
       61 gactttttcg aatgcatggc ctctttgtgg cctcccatcc ctgggaagtc atagtgggga
      121 cagtgacact gaccatctgc atgatgtcca tgaacatgtt tactggtaac aataagatct
      181 gtggttggaa ttatgaatgt ccaaagtttg aagaggatgt tttgagcagt gacattataa
241 ttctgacaat aacacgatgc atagccatcc tgtatattta cttccagttc cagaatttac
      301 gtcaacttgg atcaaaatat attttgggta ttgctggcct tttcacaatt ttctcaagtt
      361 ttgtattcag tacagttgtc attcacttct tagacaaaga attgacaggc ttgaatgaag
      421 ctttgccctt tttcctactt ttgattgacc tttccagagc aagcacatta gcaaagtttg
      481 ccctcagttc caactcacag gatgaagtaa gggaaaatat tgctcgtgga atggcaattt
      541 taggtcctac gtttaccctc gatgctcttg ttgaatgtct tgtgattgga gttggtacca
      601 tgtcaggggt acgtcagctt gaaattatgt gctgctttgg ctgcatgtca gttcttgcca
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WO 99/50454 PCT/US99/06473

42/97

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DEFINITION Human protein C gene, complete cds.
ACCESSION
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NID
            g190333
KEYWORDS
            glycoprotein; protease; protein C; serine protease.
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            The nucleotide sequence of the gene for human protein {\tt C}
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            Proc. Natl. Acad. Sci. U.S.A. 82 (14), 4673-4677 (1985)
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FIG. 23A

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LOCUS HUMLCAT 1744 bp mRNA PRT 07-JAN-1995 DEFINITION Human lecithin-cholesterol acyltransferase mRNA, complete cds, 5' and 3' flanking DNA sequences. ACCESSION M12625 NTD g187022 KEYWORDS lecithin cholesterol acyltransferase. SOURCE Human adult liver (library of A.Ullrich and L.Coussens), cDNA to mRNA, clones PL[2,4,10,12,19], and DNA. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE 1 (bases 1 to 1744) **AUTHORS** McLean, J., Fielding, C., Drayna, D., Dieplinger, H., Baer, B., Kohr, W., Henzel, W. and Lawn, R. TITLE Cloning and expression of human lecithin-cholesterol acyltransferase cDNA JOURNAL Proc. Natl. Acad. Sci. U.S.A. 83 (8), 2335-2339 (1986) MEDLINE 86205950 COMMENT Draft entry and sequence in computer readable form for [1] kindly provided by J.W.McLean, 24-JUL-1986. Because only the 5' and 3' flanking sequences were determined from DNA, it is not known whether this gene contains introns. **FEATURES** Location/Qualifiers source 1..1744 /organism="Homo sapiens" /db_xref="taxon:9606" /map="16q22.1" mRNA <257..1610 /note="LCAT mRNA" sig_peptide 268..339 /gene="LCAT" /note="lecithin-cholesterol acyltransferase signal peptide" gene 268..1590 /gene="LCAT" CDS 268..1590 /gene="LCAT" /note="lecithin-cholesterol acyltransferase precursor (EC 2.3.1.43)* /codon_start=1 /db_xref="GDB:G00-119-359" /db_xref="PID:g307117"

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       61 tgtttcccct ggcgccaaga gaagaaggcg gaactgaacc caggcccaga gccggctccc
      121 tgaggctgtg cccctttccg gcaatctctg gccacaaccc ccactggcca ggccgtccct
      181 cccactggcc ctagggcccc tcccactccc acaccagata aggacagccc agtgccgctt
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LOCUS HUMHCII 2182 bp mRNA PRI 08-NOV-1994 DEFINITION Human heparin cofactor II (HC-II) mRNA, complete cds. M12849 M19241 ACCESSION NID g183909 KEYWORDS heparin cofactor II; protease inhibitor. SOURCE Human fetal liver, cDNA to mRNA, clone lambda-HCII.7 [1]; adult liver, cDNA to mRNA, clone lambda HCII.7.1 [3]. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE 1 (bases 1025 to 2182) **AUTHORS** Inhorn, R.C. and Tollefsen, D.M. JOURNAL Unpublished (1986) (bases 1025 to 2182) REFERENCE **AUTHORS** Inhorn, R.C. and Tollefsen, D.M. TITLE Isolation and characterization of a partial cDNA clone for heparin cofactor II1 **JOURNAL** Biochem. Biophys. Res. Commun. 137 (1), 431-436 (1986) MEDLINE 86242236 (bases 1 to 2182) REFERENCE **AUTHORS** Blinder, M.A., Marasa, J.C., Reynolds, C.H., Deaven, L.L. and Tollefsen, D.M. TITLE Heparin cofactor II: cDNA sequence, chromosome localization, restriction fragment length polymorphism, and expression in Escherichia coli **JOURNAL** Biochemistry 27 (2), 752-759 (1988) MEDLINE 88163663 COMMENT [1] revises [2]. Draft entry and computer-readable sequence of [2] kindly provided by D.M.Tollefsen, 18-AUG-1986. Draft entry and computer-readable sequence of [3] kindly provided by Blinder, M.A. 24-MAR-1988. **FEATURES** Location/Qualifiers source 1..2182 /organism="Homo sapiens" /db_xref="taxon:9606" /map="22g11.2" mRNA <1..2182 /note="heparin cofactor II mRNA" sig_peptide 29..85 /gene="HCF2" /note="heparin cofactor II signal protein" 29..1528 gene /gene="HCF2" CDS 29..1528 /gene="HCF2" /note="heparin cofactor II precursor" /codon_start=1 /db_xref="GDB:G00-120-038" /db_xref="PID:g183910" /translation="MKHSLNALLIFLIITSAWGGSKGPLDQLEKGGETAQSADPQWEQ LNNKNLSMPLLPADFHKENTVTNDWIPEGEEDDDYLDLEKIFSEDDDYIDIVDSLSVS PTDSDVSAGNILQLFHGKSRIQRLNILNAKFAFNLYRVLKDQVNTFDNIF1APVGIST AMGMISLGLKGETHEQVHSILHFKDFVNASSKYEITTIHNLFRKLTHRLFRRNFGYTL RSVNDLYIQKQFPILLDFRTKVREYYFAEAQIADFSDPAFISKTNNHIMKLTKGLIKD

FIG. 25A

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08-AUG-1995

PRI

LOCUS

HUMFVA

51/97

mRNA

6893 bp

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NID
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            coagulation factor V; factor V; glycoprotein.
KEYWORDS
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            1 (bases 3636 to 6893)
REFERENCE
  AUTHORS
            Kane, W.H. and Davie, E.W.
  TITLE
            Cloning of a cDNA coding for human factor V, a blood coagulation
            factor homologous to factor VIII and ceruloplasmin
  JOURNAL
            Proc. Natl. Acad. Sci. U.S.A. 83 (18), 6800-6804 (1986)
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  MEDLINE
REFERENCE
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  AUTHORS
            Kane, W.H., Ichinose, A., Hagen, F.S. and Davie, E.W.
            Cloning of cDNAs coding for the heavy chain region and connecting region of human factor V, a blood coagulation factor with four
  TITLE
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            Biochemistry 26 (20), 6508-6514 (1987)
  JOURNAL.
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52/97

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REFERENCE
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 AUTHORS
            Wion, K.L., Kirchgessner, T.G., Lusis, A.J., Schotz, M.C. and
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            Human lipoprotein lipase complementary DNA sequence Science 235 (4796), 1638-1641 (1987)
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            Degen, S.J. and Davie, E.W.
            Nucleotide sequence of the gene for human prothrombin
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            Biochemistry 26 (19), 6165-6177 (1987)
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            Characterization of the Alu-rich 5'-flanking region of the human
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            Gene 95 (2), 253-260 (1990)
  JOURNAL
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                      3122..3415
                      /note="Alu repeat"
                      3804..4087
     repeat_region
                      /note="Alu repeat"
                      4210..4511
     repeat_region
                      /note="Alu repeat"
     repeat_region
                      4553..4793
                      /note="Alu repeat"
                      4901..5201
     repeat_region
                      /note="Alu repeat"
     protein_bind
                      4957..4962
                      /bound_moiety="Sp1"
```

```
protein_bind
                 5084..5091
                 /bound_moiety="Ap1"
repeat_region
                 5231..5443
                 /note="Alu repeat"
protein_bind
                 5231..5238
                 /bound_moiety="EBP 20"
protein_bind
                5711..5716
                 /bound_moiety="Sp1"
protein_bind
                5723..5730
                /bound_moiety="EBP 20"
protein_bind
                6047..6054
                /bound_moiety="EBP 20"
misc_feature
                6198..6237
                /note="MER sequence"
exon
                6544..6653
                /note="prothrombin precursor"
                /number=1
sig_peptide
                join(6575..6653,7040..7089).
                /gene="F2"
gene
                join(6575..6653,7040..7200,7860..7884,8127..8177,
                10504..10609,10706..10842,13181..13495,13820..13948,
                14033..14159,15317..15484,15982..16155,16698..16879,
                26327..26397,26544..26687)
                /gene="F2"
CDS
                join(6575..6653,7040..7200,7860..7884,8127..8177,
                10504..10609,10706..10842,13181..13495,13820..13948,
                14033..14159,15317..15484,15982..16155,16698..16879,
                26327..26397,26544..26687)
                /gene="F2"
                /note="precursor"
                /codon_start=1
                /product="prothrombin"
                /db_xref="PID:g339641"
```

/translation="MAHVRGLQLPGCLALAALCSLVHSQHVFLAPQQARSLLQRVRRA
NTFLEEVRKGNLERECVEETCSYEEAFEALESSTATDVFWAKYTACETARTPRDKLAA
CLEGNCAEGLGTNYRGHVNITRSGIECQLWRSRYPHKPEINSTTHPGADLQENFCRNP
DSSTTGPWCYTTDPTVRRQECSIPVCGQDQVTVAMTPRSEGSSVNLSPPLEQCVPDRG
QQYQGRLAVTTHGLPCLAWASAQAKALSKHQDFNSAVQLVENFCRNPDGDEEGVWCYV
AGKPGDFGYCDLNYCEEAVEEETGDGLDEDSDRAIEGRTATSEYQTFFNPRTFGSGEA
DCGLRPLFEKKSLEDKTERELLESYIDGRIVEGSDAEIGMSPWQVMLFRKSPQELLCG
ASLISDRWVLTAAHCLLYPPWDKNFTENDLLVRIGKHSRTRYERNIEKISMLEKIYIH
PRYNWRENLDRDIALMKLKKPVAFSDYIHPVCLPDRETAASLLQAGYKGRVTGWGNLK

ETWTANVGKGQPSVLQVVNLPIVERPVCKDSTRIRITDNMFCAGYKPDEGKRGDACEG

```
DSGGPFVMKSPFNNRWYQMGIVSWGEGCDRDGKYGFYTHVFRLKKWIQKVIDQFGE*

6654..7039
/note="prothrombin intron A"

7040..7200
/gene="F2"
/number=2
join(7090..7200,7860..7884,8127..8177,10504..10609,
10706..10842,13181..13495,13820..13948,14033..14159,
15317..15484,15982..16155,16698..16879,26327..26397,
26544..26684)
/gene="F2"
/product="thrombin"
```

FIG. 28B

```
intron
                  7201..7859
                  /note="prothrombin intron B"
 exon
                  7860..7884
                  /gene="F2"
                  /number=3
intron
                  7885..8126
                  /note="prothrombin intron C"
 exon
                  8127..8177
                  /gene="F2"
                  /number=4
intron
                  8178..10503
                  /note="prothrombin intron D"
repeat_region
                  8330..8675
                  /note="Alu repeat copy A"
repeat_region
                  9030..9161
                  /note="Alu repeat copy B"
repeat_region
                  9176..9475
                  /note="Alu repeat copy C"
repeat_region
                  9643..9937
                  /note="Alu repeat copy D"
exon
                  10504..10609
                  /gene="F2"
                  /number=5
intron
                  10610..10705
                  /note="prothrombin intron E"
exon
                  10706..10842
                  /gene="F2"
                  /number=6
variation
                  10774
                  /gene="F2"
                  /note="c in DNA; a in cDNA"
intron
                  10843..13180
                 /note="prothrombin intron F* 10933..11232
repeat_region
                  /note="Alu repeat copy E"
repeat_region
                 12089..12390
                  /note="Alu repeat copy F"
repeat_region
                 12391..12689
                 /note="Alu repeat copy G" 13181..13495
exon
                  /gene="F2"
                  /number=7
intron
                 13496..13819
                 /note="prothrombin intron G" 13820..13948
exon
                 /gene="F2"
                 /number=8
intron
                 13949..14032
                 /note="prothrombin intron H"
exon
                 14033..14159
                 /gene="F2"
                 /number=9
intron
                 14160..15316
                 /note="prothrombin intron I"
repeat_region
                 14325..14643
                 /note="Alu repeat copy H"
repeat_region
                 14820..15126
                 /note="Alu repeat copy I"
exon
                 15317..15484
                 /gene="F2"
                 /number=10
intron
                 15485..15981
                 /note="prothrombin intron J"
exon
                 15982..16155
                 /gene="F2"
```

FIG. 28C

```
/number=11
                 16156..16697
intron
                 /note="prothrombin intron K"
repeat_region
                 16306..16596
                 /note="Alu repeat copy J"
exon
                 16698..16879
                 /gene="F2"
                 /number=12
intron
                 16880..26326
                 /note="prothrombin intron L (no splice consensus at
                 16880); putative*
repeat_region
                 16952..17098
                 /note="potential new repetitive element copy A; putative"
17145..17206
repeat_region
                 /note="potential new repetitive element copy B; putative"
                 17375..17614
repeat_region
                 /note="Alu repeat copy K"
                 18250..18531
repeat_region
                 /note="Alu repeat copy L"
                 18545..18795
repeat_region
                 /note="Alu repeat copy M"
                 19231..19527
repeat_region
                 /note="Alu repeat copy N"
                 19706..20012
repeat_region
                 /note="Alu repeat copy 0"
                 20584..20815
repeat_region
                 /note="Alu repeat copy P"
                 21088..21375
repeat region
                 /note="Alu repeat copy Q"
                 21120..21290
repeat_region
                 /note="KpnI repeat copy A"
21387..21539
repeat_region
                 /note="Alu repeat copy R"
repeat_region
                 21814..22110
                 /note="Alu repeat copy S"
                 22315..22434
repeat_region
                 /note="Alu repeat copy T"
                 22441..22738
repeat_region
                 /note="Alu repeat copy U"
repeat_region
                 22748..22921
                 /note="Alu repeat copy V"
                 22922..23203
repeat_region
                 /note="Alu repeat copy W"
                 23204..23496
repeat_region
                 /note="Alu repeat copy X"
                 23558..23876
repeat_region
                  /note="Alu repeat copy Y"
repeat_region
                 24037..24363
                  /note="KpnI repeat copy B"
                 24421..24720
repeat_region
                  /note="Alu repeat copy Z"
                 24721..25015
repeat_region
                  /note="Alu repeat copy AA"
repeat_region
                  25112..25282
                  /note="Alu repeat copy AB"
                  25283..25575
 repeat_region
                  /note="Alu repeat copy AC"
                  25752..25998
 repeat_region
                  /note="Alu repeat copy AD"
                  26327..26397
 exon
                  /gene="F2"
                  /number=13
 intron
                  26398..26543
                  /note="prothrombin intron M"
```

```
exon
                          26544..>26687
                          /gene="F2"
                          /note="prothrombin precursor"
                          /number=14
      polyA_signal
                          26765..26770
      repeat_region
                          26881..26928
                          /note="Alu repeat copy AE"
BASE COUNT
                  6463 a
                             6624 c
                                                    7086 t
                                        6755 g
ORIGIN
         1 gcgtgagcca ctgcgccctg accacatata atttttatta attataatgt tgaaagtccc
        61 tttattccac accteteete teatteacte etggtaggte atttttaatg atttgatgta
       121 tatactgaat ttggatgctt cttgctacag ggcaaagacg ctaataagat tttgctggag
       181 ccttttcaca gatgcaagtc aatccaggca gtgtctatag ctgctgaacc caaaatcaga
       241 aagcgagggc tatcaaagct cttctgtcct gatttgcaac tttagtagtg caagaaaaaa
       301 aatettagaa taaaaaatgg gtacegttca gagacettta gagattgcaa ggcatcacag
       361 atgataaaaa gctccatctc tagacgtgtt caggagtggg ttggggcttt gaccttgact
       421 agetgeatea aettggacaa gteaettege tteeetgtge etcagtttee teatecataa
       481 aatggggata agtatagtac ctacctcata agtcctgcct acctagcaca tggtgagcaa
       541 ttactaaatt gtaggcctag teectataat eccageaett ttggagaaca aggtagggga
       601 atcgcttgaa gccaggagtt ccagaccagc ctggccaaca tagtgagact gtgtttctat
       661 aaaataaaaa aaaaaaatac ccaagcttgg tggtgcaggc ctgtagtccc ggctacttgg
       721 gagtetgagt caggaggatt gettgagece aggagtteaa ggttgtagta agetatgatt
      781 gcaccactgc actccagcct ggcgacagag catgaccctg tctctaaaaa tataaaatta
841 ggccaggcac agtggttcat gcctgtaatt ccaacatttt gggaggccaa ggcaggtgga
      901 tcactgtgag ctcagcagtt cgagaccagc ctgggcaaca aggcaaaatc ctgtctctac
     961 taaaattaca aaaattagcc aggagaggtg gtacacgcct gtaatcccag ttactgggga
1021 agctgaagca ggagaattgc ttgaacccgg gaggcgaagg ttgcagtgag ccaagatcgt
     1081 gccattgcac tgcagcctag gagacagagc gagactcgat ctcaataaat aaataaatta
     1141 attaattaat aaaaaaataa gttgggcatg gtggcacctg cctgtagtcc aagctactca
1201 ggaggctaga ggtgggagga tcacttgagc caggagttct aggctgcagt gagctattat
     1261 cacgccacca tactccagcc tgctgtatgt actccagcct gggcaacaga gtgacaccct
     1321 gtctcaaagt aaagtaaaat aaaaattaaa aaacaaatta ctaaattgta cttaacagta
     1381 ttgtcatcag tcttcctaaa taggaggaca ggcaaaatta agggacttaa catgtgccct
     1441 caggtatagt agtttggggc aggccagcat cacccgcaca gtagttctgt actgtaggtg
     1501 cgtgttctct gggtcaactt tatggcccag tgaggccgta ctctaccaga atgtcagggg
     1561 acaagggttg ggagaggcaa aagtgctggt ctgaagcagg agtctgggtt tccatcctag
     1621 ctctaccacc aattctgtat gaccgtgccc cctccatttc ctccatgacc acatagagac
1681 atggggcagt tggatgaaat caatgattcc cagtcttggc tctatcatgg aaccatttgc
     1741 taacttettt ttttetetta tggateecat atttttaaag atttttaeta aatagaaatt
     1801 gacttatact tttccaagct ggagtgtggt ggcatgattt cagctcactg caacctccgc
     1861 ctcccgggtt caagtgattc tcctgcctca gcctcctgag tagctgggat tataggtgct
     1921 caccaggccc ggctaattt tttgtattt tagtagagac agaatttcac catgttggcc
1981 aggctgattt caaactcctg acctcaagtg atctgctcac ctcagcctcc caaagtgctg
2041 ggattacagg cgtgagtcac tatgcccagc cgcttactca cattttctag tcaaaataga
     2101 aaactgctta agtcactgtc tgcagaagag caaaaaaaaa aaaagaaata aaaaattgaa
     2161 aactgctgat cagattgaga aaaacataag attattcacc acctaaagag aaaaaatttc 2221 agtcgaaagg gaaaaaaatt catttttgtc ttaataaggc aaattcacaa tttttgaggt
     2281 tttaacaaaa tatatgcaga aagacaaggc cacccgtag aacgtgcaca cagccctagg
     2341 cttggaaatg gctggattta ataatatctg gtctttcttt gagccctgaa attctctaac
2401 actatgtctt ggaacataat tttactgttt tcagtggtta tagagatttg ctttacaatt
     2461 tagcattggt ctttacccat gattttgttt gacgccaact tgttggcagg aatgcacccc
    2521 ctgcccccg ctttgttatg gccttgctcc tatagggcaa gaatatctgc tttaaggccg
    2581 ggtgtggtgg ctcaggcctg taatcccagc actttgaggg gccaaggcgg gcagatcacc
2641 tgaggtcagg agtttgagac cagcctggcc agtatggtga aatcctgtct ctactaaaaa
    2701 taacaaaaat tagctgggtg tggtggcaca cacctgtaat cccagctatt tgggaggccg
    2761 aaacaagaga accacttgaa cccaggaggc ggaggttgcg gtgagccgag attatgccac
2821 tgcactccag cctgggaaac agagcaagat tccgtctcac acacaaaaaa tatatatatg
    2881 tctgctttaa gtatgcaggc cgtgtttgtg ctgaacggca ggaatgccaa acttggctgc
    2941 atggtaccaa ctagggacct cagagttcca aggagaacaa acagttggtt cctggaggct
    3001 gggggcttgt atcagaccct gaagactaag catgtgctgg gtccattgtt gtcctgcacc
    3061 catggtagtg cactaaacac ctaacctata tttaagtgtt tttgtttgtc caaaaaatgt
    3121 ctttttttt tgggagtcaa gagtcttgct ctgttgccca ggctggagtg cagtgacacg
    3181 atctcagctc actgcagcct ccgcctcccg ggttcaagct attctcctgt ctcagcctcc 3241 caaatagctg agactatagg cacgcacatc catgcccagc taatttttt atttttagta
```

```
3301 gagacgaggt gtctccatgg tggccaggtt ggtcttgaac tcctgtcctc aagtgatcca
 3361 cctgcctcgg cctcccaaag tggtgggatt gcaggcatga gacaccgcgc ccggcctgcc
 3421 ttgtcccttc ttaaaatgag ttgtccattt gtaagctgct gatttctttg ggacattgtc
 3481 teegtaaact tttcataaag catcagtgat ttcaccatte ttccacccaa gettcaccgt
 3541 aaatttgttg tttgttcttg cttcaatttc agcagaattc atttagctct gataagggct
 3601 cgcttcaaac tgatgtctta tccttcttag tgcctcaaac tacatcctgt tcactcatgt
 3661 tatagcaagt tagtgtgagt ttattttggt gcacaaaaat ttttttaaat ccatgcagtc
 3721 ttttttcata atacgcattt tccatgaact tttcgaagac cccttgtaga tgtctgttgt
 3781 ttaaaccacc cagtttacag taatttttt tttttttga gatgaagtct tgctctgtcg
 3841 cccaggctgg agtgcattgg cacactctcg gctcactgca acctctgcct cctgggttca 3901 agcaattttt ctgtctcagt ctcccgagta gctgggatta caggtgtgtg ccaccatgcc
 3961 tagctaattt atgtgttttt agtagagacg gggtttcact atgttggcta ggctggtctc
 4021 gaacteetea cettgtgate ggeeegeete ggeeteecaa agtattggga ttacaggegt
4081 gagactettg caettggeet acagtaattt tatagcagee taggetaaga tagccattte
 4141 tgggtataag aatgtcatat actgaacagg cctgcaactg tgagtaaaag tctgcaaaga
 4201 ggccgggcag tggctcatac ctgtaatecc ageaetttgg ggggccgagg caggtggate
 4261 acctgaggtc agcagttcga gaccagcctg accaacatgg tgaaacccca tctctactaa
 4321 aaatacaaaa ttagctgggc gtggtagtgc atgcttgtaa tccctagcat gcacttggga
 4381 gctacttggg aggctgaggc aggagaatca cttgtactca ggaggccgag gttgcagtga
 4441 getgagatea egecaetgea eteettetg ggtgacagag tgagaeteea teteaaaaaa
 4501 acaaaacaaa acaaacaaa aacaaaccaaa aggtaggtag cagtggttca
4561 cgcctgtaat ccccactttg gaggctaaag tgggcagatc acctgaggtc aggagttcac 4621 gtccagcctg ggcaacatgg tgaaactctg tcctacaaa aatacaaaaa ttagccaggc 4681 atgatggcgg gtgctgtagt tccagctatt cgggaggetg aggcagagg atcgcttgaa 4741 cctaggaggt agaggttgca gtgagccgag ttcacgctat tgcactccag cctccatctc
4861 tttattttat tttatttttc taggaacagg tctcattcag gccaggcatg gtgctcacgc
4921 ctgtaatccc agcacttggg aggccgaggt ggaggtgggc ggatcacctg aggtcaggag
4981 ttcgagccat cctggtcaat gtggcgaaac cccatctcta ctaaaaatac aaaaattagc
5041 caggtgtggt ggcacacgcc tgtaattcca gctacttggg atactgagtc aggagaatca
5101 cttgaacagg gagatggaaa ttgcagtgag ccgagattgt tccactgcac tccagcctgg
5221 agaaacagga tettactetg ttacccagge tggagtacag tggtgcaate atageteact
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5341 tgctatgttg cccaggctgg tctcaaactc ctggcctcaa gcgatcctgc catgtcggcc
5401 teccaaagtg ttgggattae aagtgtgage cactatgeet ggeetaaaaa tatatatatg
5461 aaaatatata agaaatggge etcecaggaa ttaaggtgtt tgegggagte etggteeca
5521 gtttttctgc caacactccc tgttcccaca catgacctgg tccagacccc aaacagccag
5581 gcccaaagga caggtgaggc gaggcgagaa cttgtgcctc cccgtgttcc tgctctttgt
5641 ccctctgtcc tacttagact aatatttgcc ttgggtactg caaacaggaa atgggggagg
5701 gacaggagta gggcggaggg tagggtagga ccagaagcct ctctaggcct gccatggggc
5761 aggcagccag ggagaaggag ggcccctcag tggagaccca gggatttcag tagcccctgt
5821 tccgggacag gcgcaggtcc tgggaggtga cagaagatag actaaaggcc caagagtccc
5881 tggacctgac tcctcccagc agctgccaca cacaaacaca cctccaggca ccctggacag
5941 gaaggaggag aaatgggccc ctcctccagt ggctgagaag ctggggcaaa tgttggctgt
6001 toctatocot ggtgcatoco atggcgaggg gcaacttoca toaggccaca cottttatot
6061 ttgtototat ttttgatato tgtgtattat gattatacaa accoccacat tggcctatat
6121 gtgcagatct gattaagaac ttacgatatt ccatggacat tccattccta atctccttta
6181 gtcctcacaa caaagtatta ttcccattgt atagatgagg aaactgaggc acacagagat
6241 gacaagcaac caccgctata tgttaggatt cgaaggagct ccaggaaagt ctcatagccc
6301 cactggccag aatgggctaa atctcagagg gggagggtgg gagatggggg tgacagtgac
6361 cttttttgtg actcctccta gaccatccat ccctgctccc aggaggacct gtcctcccag
6421 atggtggaga tggacaggag gactatctac ccacccgtcc ccacggccct gaccctctga
6481 ceteacete teegetgatt tetteatgtt agtteaacat tacceagagg ggteaggaca
6541 gacaatteet cagtgaceca ggagetgaca caetatggeg caegteegag gettgeaget
6601 gcctggctgc ctggccctgg ctgccctgtg tagccttgtg cacagccagc atggtaaggg
6661 agtgcttgca ggctggaaca ggctggagga ctggggtgtg ggcccatggg ctggggtctc
6721 ctggctggac agagcacaca gagctggccc ctaagtaggt ctcagcccca ggcggccagc
6781 ttagggaaga agtcaggagc tcagggctgg aaagagaatg gctgcttctc tcttccaata
6841 tagggagcag gctgggggca aggggcagtg taggaggggc acagggggcc acatttagca
6901 gccttccagg ccttccacca gcccagactg cctctctcag aagccagcag gggagggtgg 6961 gcttgcttca tgcccccaga tggccaagac tgcctgttcc tgaggtcgct gttccatgac
7021 ccccccaccg cctttacagt gttcctggct cctcagcaag cacggtcgct gctccagcgg
7081 gtccggcgag ccaacacctt cttggaggag gtgcgcaagg gcaacctaga gcgagagtgc
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  7201 gtgagcctgg gctgctcgga cggtgccggg gcctcagacc gggcccaact ctagacactt
  7261 ccacagagaa gcaagcgagg aacgccacag cccttcgct gctcacagcc tcatttcaac
  7321 tetgageece teetcacagg getggcaaga ggageggeet cageetttee tgggggtete 7381 tgtgeetgga etgtgteect gtgcagetee atgaeatggg gaggeeteea cagtetteag
  7441 acatecacet geettggage tetgtgteca catggeetee teageggeag acteceacae
  7501 caccettgag gggtgggact ctggggaggc caccacaagc ccccgggctc aagactcagt
  7561 gttcctggag ctctgtgtcg cctttcctgt ctgtagggac tctgccaggg acccactgcc
  7621 ccctctcctc ccatctcccc cagcctcttt cagactcggt gtgtgtgttg gaggaactcc
  7681 cctatcctca aatattcttc tccttttgga aacaaaagta ggaaactctg ccacaaacct
  7741 ccccagagcc tgccccctgc gtgaccaggg taaggaaagt gtgaggagga gcataacatt
  7801 tactaaaaca acacaaaaca ggagctgccg tagcctcact cccagccctt gtttttcagg
  7861 atgtgttctg ggccaagtac acaggtgagc accgggaagg atttgcccca ggaagggagg
  7921 cctggggacc ccagtgagag aattctaccc agagaatctt ctgctgcacc tagccatcca
  7981 cccatccacc cettececac tecttecttg gtecetecca tetgtteate catettetg
  8041 tttctcacca acateccate caceetgact ccageteate etggecatae eccaatecca
  8101 aaggtaaaca cetgggtett ttecagettg tgagacageg aggacgeete gagataaget
  8161 tgctgcatgt ctggaaggtg agcaactgac acgggtttgg ggagcaggac atggagggga
  8221 gcttgggaga agagctcagg ggtgggtttg gagtgtggct ggtggaggcc gaggcagtcc
  8281 ccagcatetg acattgetee catteetggg gtcaagatgt etetttgtae etggetetgt
  8341 gtctggcatg cgaacgaatg aatgaatgaa tggactaatg aattaatgtt ttttttttg
  8401 agacagagtc tcgctctgtt gcccaggctg gagtgcagtg gcacgatctt ggctcactgt
  8461 aaactccgcc tcccggattc aagcaattct ctgcctcaac ctcccaagta gctgggatta
8521 caggtgctcg ccaccacgcc tagctaattt ttgtattttt agtagagacg gggtttcacc
 8581 atgttggcca ggctggtctt gaactcctga cctcgtgatc cacccacctc ggcctcaaag
 8641 tgctgggatt atagaagtga gccaccgcgc ctggccatga attcatgttt aaggcttcat
  8701 teteetttge etgaccegag tetetgecce cacetagtea gagetttgat gatgteacat
 8761 teceetteta getttaggtg teaetgaace aaacaggaac ccaaaceccc agetgetetg
 8821 acaccaagga cttccctaag catgccaagg tgtttctagc acctggcctt gcatatgttg
8881 tcaatttcct ctggagcgac catcacatct actgaacact ttcctatcct tcaaggactg
 8941 cttcaaatgt caccactttt gctgagactt cagggagcac cctccctcct gcactgtgtc
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20/01	tcccaataaa	agtgactctc	agcgagcctc	aatgctccca	gtgctattca	tgggcagete
26821	tctgggctca	ggaagagcca	gtaatactac	togataaaga	agacttaaga	atccaccacc
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LOCTIC HUMPMG3BA 3997 bp mRNA PRI 08-JAN-1995 DEFINITION Human platelet membrane glycoprotein IIIa beta subunit mRNA, complete cds. ACCESSION M20311 g190107 NID KEYWORDS cell membrane glycoprotein; platelet membrane glycoprotein IIIa. SOURCE Homo sapiens cDNA to mRNA. ORGANISM Homo sapiens Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo. 1 (bases 1 to 3997) REFERENCE AUTHORS Zimrin, A.B., Eisman, R., Vilaire, G., Schwartz, E., Bennett, J.S. and Poncz, M. TITLE Structure of platelet glycoprotein IIIa. A common subunit for two different membrane receptors J. Clin. Invest. 81 (5), 1470-1475 (1988) JOURNAL 88213696 MEDLINE FEATURES Location/Qualifiers source 1..3997 /organism="Homo sapiens" /db_xref="taxon:9606" /cell_type="erythroleukemia" /map="17q21.32" sig_peptide 17..94 /gene="ITGB3" /note="G00-120-013" CDS 17..2383 /gene="ITGB3" /codon_start=1 /db_xref="GDB:G00-120-013" /product="glycoprotein IIIa" /db_xref="PID:g190108" /translation="MRARPRPRPLWATVLALGALAGVGVGGPNICTTRGVSSCOOCLA VSPMCAWCSDEALPLGSPRCDLKENLLKDNCAPESIEFPVSEARVLEDRPLSDKGSGD SSQVTQVSPQRIALRLRPDDSKNFSIQVRQVEDYPVDIYYLMDLSYSMKDDLWSIQNL GTKLATOMRKLTSNLRIGFGAFVDKPVSPYMYISPPEALENPCYDMKTTCLPMFGYKH VLTLTDQVTRFNEEVKKQSVSRNRDAPEGGFDAIMQATVCDEKIGWRNDASHLLVFTT DAKTHIALDGRLAGIVQPNDGQCHVGSDNHYSASTTMDYPSLGLMTEKLSQKNINLIF AVTENVVNLYQNYSELIPGTTVGVLSMDSSNVLQLIVDAYGKIRSKVELEVRDLPEEL SLSFNATCLNNEVIPGLKSCMGLKIGDTVSFSIEAKVRGCPQEKEKSFTIKPVGFKDS LIVQVTFDCDCACQAQAEPNSHRCNNGNGTFECGVCRCGPGWLGSQCECSEEDYRPSQ QDECSPREGQPVCSQRGECLCGQCVCHSSDFGKITGKYCECDDFSCVRYKGEMCSGHG QCSCGDCLCDSDWTGYYCNCTTRTDTCMSSNGLLCSGRGKCECGSCVCIQPGSYGDTC EKCPTCPDACTFKKECVECKKFDREPYMTENTCNRYCRDEIESVKELKDTGKDAVNCT YKNEDDCVVRFQYYEDSSGKSILYVVEEPECPKGPDILVVLLSVMGAILLIGLAALLI

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121 tgtgagctcc tgccagcagt gcctggctgt gagccccatg tgtgcctggt gctctgatga
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LOCUS
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                                                                     31-OCT-1994
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ACCESSION
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NID
             antithrombin; antithrombin III.
KEYWORDS
SEGMENT
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SOURCE
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             Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
             1
                (bases 1 to 238)
             Bock, S.C., Marrinan, J.A. and Radziejewska, E.
  AUTHORS
             Antithrombin III Utah: proline-407 to leucine mutation in a highly
  TITLE
             conserved region near the inhibitor reactive site [published
             erratum appears in Biochemistry 1989 Apr 18;28(8):3628]
Biochemistry 27 (16), 6171-6178 (1988)
  JOURNAL
  MEDLINE
             89050967
COMMENT
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BASE COUNT
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121 agagaagttc ctctgaacac tattatcttc atgggcagag tagccaaccc ttgtgttaag
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                                                      PRI
                                                                08-NOV-1994
DEFINITION
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ACCESSION
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NID
            g183449
KEYWORDS
            platelet glycoprotein IIb.
SEGMENT
            2 of 2
SOURCE
            Homo sapiens (tissue library: lambda-EMBL 4) DNA.
  ORGANISM
            Homo sapiens
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            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 1 to 623)
  AUTHORS
            Prandini, M.H., Denarier, E., Frachet, P., Uzan, G. and Marguerie, G.
            Isolation of the human platelet glycoprotein IIb gene and
  TITLE
            characterization of the 5' flanking region
            Biochem. Biophys. Res. Commun. 156 (1), 595-601 (1988)
  JOURNAL.
  MEDLINE
            89025907
COMMENT
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      121 acttecteae atgtgetetg gggeeageaa ateatetgta taccetgace ttggeeeeeg
      181 tgtaccccca ggtcggcttc ttcaagcgga accggcacac cctggaagaa gatgatgaag
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75/97

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LOCUS
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                                                                01-NOV-1994
DEFINITION Human cholesteryl ester transfer protein mRNA, complete cds.
ACCESSION
            M30185
            g180259
NID
KEYWORDS
            cholesteryl ester transfer protein; transfer protein.
            Human adult liver, cDNA to mRNA.
SOURCE
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
            1 (bases 1 to 1787)
 AUTHORS
            Drayna, D., Jarnagin, A.S., McLean, J., Henzel, W., Kohr, W.,
            Fielding, C. and Lawn, R.
 TITLE
            Cloning and sequencing of human cholesteryl ester transfer protein
            CDNA
  JOURNAL
            Nature 327 (6123), 632-634 (1987)
 MEDLINE
            87258172
FEATURES
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     1201 caattettea gtgatggtga aatteetett teeaegeeca gaccagcaac attetgtage
     1261 ttacacattt gaagaggata tcgtgactac cgtccaggcc tcctattcta agaaaaagct
     1321 cttcttaagc ctcttggatt tccagattac accaaagact gtttccaact tgactgagag
     1381 cageteegag tecateeaga getteetgea gteaatgate accgetgtgg geateeetga
     1441 ggtcatgtct cggctcgagg tagtgtttac agccctcatg aacagcaaag gcgtgagcct
     1501 cttcgacatc atcaaccctg agattatcac tcgagatggc ttcctgctgc tgcagatgga
     1561 ctttggcttc cctgagcacc tgctggtgga tttcctccag agcttgagct agaagtctcc
     1621 aaggaggteg ggatgggget tgtageagaa ggeaageace aggeteaeag etggaaeeet
1681 ggtgteteet eeagegtggt ggaagttggg ttaggagtae ggagatggag attggeteee
     1741 aactcctccc tatcctaaag gcccactggc attaaagtgc tgtatcc
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LOCUS
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                         13204 bp
                                      DNA
                                                       PRI
                                                                  10-NOV-1994
DEFINITION
            Human platelet Glycoprotein IIb (GPIIb) gene, exons 2-29.
ACCESSION
            M33320
            g183506
NID
            platelet Glycoprotein IIb.
KEYWORDS
SEGMENT
            2 of 3
SOURCE
            Human leukocyte DNA.
  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
                (bases 1 to 13204)
  AUTHORS
            Heidenreich, R., Eisman, R., Surrey, S., Delgrosso, K., Bennett, J.S.,
            Schwartz, E. and Poncz, M.
            Organization of the gene for platelet glycoprotein IIb Biochemistry 29 (5), 1232-1244 (1990)
  TITLE
  TOURNAL.
  MEDLINE
            90212612
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     exon
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     exon
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                      /note="GPIIb intron E (no splice consensus); putative;
                      does not fit consensus*
     exon
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                      1552..1680
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     exon
                      2042..2089
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/number=8 intron 2090..2244 /note="GPIIb intron H (no splice consensus); putative; does not fit consensus" exon 2245..2288 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=9 2289..2460 intron /note="GPIIb intron I" 2461..2514 exon /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=10 intron 2515..2652 /note="GPIIb intron J" exon 2653..2705 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=11 intron 2706..2896 /note="GPIIb intron K" exon 2897..3108 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=12 intron 3109..5535 /note="GPIIb intron L" exon 5536..5718 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=13 intron 5719..5951 /note="GPIIb intron M" 5952..5997 exon /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=14 intron 5998..6105 /note="GPIIb intron N" exon 6106..6210 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=15 intron 6211..6294 /note="GPIIb intron 0" exon 6295..6350 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=16 intron 6351..6442 /note=*GPIIb intron P* exon 6443..6594 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=17 intron 6595..6782 /note="GPIIb intron Q" exon 6783..6908 /gene="ITGA2B" /note="platelet Glycoprotein IIb" /number=18 intron 6909..7885 /note="GPIIb intron R" exon 7886..7953

FIG. 33B

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                          3579 с
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                                               2722 t
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       61 agaggtaccc gctaccttcc ctcattaaaa ccagctctca agaggggatc tggtaacagt
      121 ctaggcaggc attccaggga gcatgtgaac cgctggttct tgttgcgggt ggaggatgga
      181 ggtgttgtac agagtttagg tcttttcag caaagatctc caaaccccgg gtgttcaaaa
      241 tcaaaccaaa ggggattata gtcccagctc tactcacaac tcactggtta ctttagccac
      301 gagattgccc tcgctgagag tcggtttcac tgtccataag atgaagaagt acatcacggt
      361 ggtctgtgag gtgtcattga ggaaagatgg tccagtgccc ccatgccaca tggccttcgg
      421 gcagtgctcc cagcgccggc gccagggcct gggatacgct ggaatctgcg cggcgctcac
      481 ccagetttee tatgeagagt ggccategtg gtgggegeec egeggaeect gggeeceage
      541 caggaggaga cgggcggcgt gttcctgtgc ccctggaggg ccgagggcgg ccagtgcccc
      601 togotgotot ttgacotoog tgagtoccag gcaaggagag caaggttggg gtcagaggga
      661 cgtggactgc ccgggcttca gcgccccacc ccttcttgtg ccttccaggt gatgagaccc 721 gaaatgtagg ctcccaaact ttacaaacct tcaaggcccg ccaaggactg ggggcgtcgg
      781 tcgtcagctg gagcgacgtc attgtggtgg gccccgcggt acagggcaca gggaacaatc
      841 gggggcaggg acactggggc caggaggagc ccaagteteg egeceegtee ccatetgtgg
      901 ccetttetca ggcetgegee ceetggeage actggaaegt ectagaaaag actgaggagg
      1021 agtactcccc ctgtcgcggg aacaccctga gccgcattta cgtggaaaat gattttagta
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     3661 ctgtgaaata agaggcccag gatagagccc tagggagcaa aagcatttag gtgactccta
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            g183907
NID
KEYWORDS
            heparin cofactor II; serpin.
SOURCE
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  ORGANISM
            Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
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REFERENCE
                (bases 1 to 15849)
  AUTHORS
            Herzog, R., Lutz, S., Blin, N., Marasa, J.C., Blinder, M.A. and
            Tollefsen, D.M.
  TITLE
            Complete nucleotide sequence of the gene for human heparin
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            II and mapping to chromosomal band 22q11
  JOURNAL
            Biochemistry 30 (5), 1350-1357 (1991)
  MEDLINE
            91120782
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FIG. 34A

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5941 caggagtttg agaccagcct ggccaacaga gcaaaacccc atctctacta aacaaaatac
6001 aaaaattaac caggcgtagt ggtgtgcacc tgtagtccca gctacttggg aggctgaggc
6061 aagagaattg cttgagtcca ggaggccgaa gttgcagtaa gccgagatcg cgccactgca
6121 ctccagcctg ggtgacagag tgagactcca tttcaaaaaa taaaaacaac aaaagccaat
6181 tacaacaaca acaacaaaaa aacaacgaat taaacaaccc caaagattgc acaaatttca
6241 agtatettta gaatatgttt teagaaagee tggeecatgg acatttttea acageatete
6301 cattgcaaag gtggaatggt gtgagtcaca caggcatggc tgagtcccac taatgcacat
6361 cccttctagg tactctccaa tcaccagccc caggtgccca ctcaagccca gctcttagtg
6421 aggtttccct gactctctgg gcacttccac tcctaccaca cagggtagag ccacacccct
6481 ttccgtaccc ccatgtgctc tggcagcatt attttgagag ccttcgcttt actgcacgtc
6541 tgtcccatct gtcccctgac tggtccatga gcccctggtg ggaactttgt ctctggtaac
6601 taaacactgt ctggaggtgg tggacaaggt gtctggagaa aaacaaactc ctccctggga
6661 tgcctgagct cccaggattc tagaaggtta gttttgcaaa cctttaaaga agggattttc
6721 atcaaggggc ccacagatcc ttcattgagg tttatgagtc ccacatcaaa ggttgggtgt
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					:	
6781	l ctatctacat	cagattctct	taaagtccar	gatcctaaaa	Carttangan	ataataataa
	- 949996666	uuulla	Laduccacado	. <u> </u>	+ ~+ ~~~ + ~ ~ +	
	- 5099466496	Cayyccucc	. Lucacioror			_
		uvilaciaaa	L CUCACI FFF	. affffcctos	****	
7081	ccccagtaga	agcagttaaa	taacaaaaa	ctgagcatgc	aaactgctca	gtctgcagat
7141	cacaaggaaa	acaccotcac	Caacaaaaac	attccagagg	CTCTTCTCCC	tgccgacttc
7261	tcagtttccc	CGacagactc	tastatasat	gctgggaaca	tcgacatcgt	cgacagtctg
7321	. aagagccgga	tccagcgtct	taacatooto	geegggaaca	tcctccagct	ttttcatggc
7381	. gtgctgaaag	accadatcaa	Cacttteent	aacgccaagt	tcgctttcaa	cctctaccga
7441	. actocoatoo	gtatgatttc	cttaggtata	aacatcttca	tagcacccgt	tggcatttct
7501	attttgcarr	ttaaacactt	tattaataa	aagggagaga	cccatgaaca	agtgcactcg
7621	tcagtcaatg	acctttatat	ccategeete	ttcaggagga	attttgggta	cacactgcgg
		CCUCCAACAC	aluadoraaa	ACCOPTACCO	ACCOMPANDS.	
		cygaaqucau	auallucant	TATCCCTSCS.	+~+~~~~	
		auctactagu	uuucaaaacc	2224644444		
	-3-3-0		CCCdarrarc	2212200200	~~+~+~	
		cegaaccacc	uuucaaaraa	CSECOCOSSS	+	***
		ggccaaacac	aauuucacer	LLLSSWWSWW	+02000000	
		CCCCCGGGG	CCLAUDAAFA	aaraammatm	~~+++	
		u caccccac	LLUUACCAGA	ccanaanaaa	~~+~+	
			Caccaaooac	Taccadataa	ataa	
	3	CCACCCCCCC	CCLLOOPINA	TTTCCCCT TTC	antatana.	
	-599449	gccaaacada	ULCACECEGA	Cacccacce	+	
		accacacaca	Calualdaac	acacatataa	+++++	
10441	cacaaactat	tatgtggccc	aaatotcaaa	aatgctgagg	cacagggcag	cccacattgc
0621	ccttgaacaa	gttacttcac	ttetttetee	ctctgtttcc	CLCTacccac	cagctagaaa
	<u> </u>				ccatatgtaa	aagagggata

					;	
1068	l acaaaacgca	a cacaacttgo	atgttgctag	r gagcagaaat	· cacataatac	aggaaaggtg
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10983	l ctcatgccta	a taatcccage	gatgaaggee	accaagacag	agctgaggct	ggcagggtgg
11701	. tttccatgat	gcagaccaag	gggaacttcc	traragraga	taaccaccac	stagicaagg
11941	ggggtgtctg	ggaatactgg	aaaataaata	grattate	ttttgagete	ccagatgctg
12001	gtacccaaga	ggaatactgg	addatygatt	accectaa	aaagggagaa	ttatgtacaa
12181	gcttcatcat	gcagatgact	Lagagacage	ttggtgcttg	ctttgtggct	tcgagtccca
12721		tttttttga	gacagggtct	tgttctgtca	cccaggctgg	agtgcagtgg
12901	tectectgee	ttggccttcc	aaagtgctgg	gattaacagg	cataaaccac	tatacctaac
13021	ccatttgact	tttaattgag	atcttacttg	gtgcaaggta	tgagctaggt	agagaagga
13441	tgcaacagaa	aacacacctc	agttttcagt	accomments	cagececaag	taattaatta
13561	aaatataacc	cgtggccctt	taaagggaaa	atcatcatta	ttttttt	accetgegat
13681	caagetggag	aagaactaca	atctagtgge	cayaacccga	gaagtgette	tgccgaaatt
13741	gtttgacaaa	aatggcaaca	toocagegga.	gtccctgaag	ttgatgggga	tcaggatgct
13861	ccacttaccc	ttcctaccca	CCCCCCCC	cccagggtet	gcctcagcac	agccccacct
13921	tetteggeet	gggtgggata agacacttac	Cacacata	catgtccca	gcttggggtg	ctgagtctgc
13981	ggcacctggc	acacacttac	tacayaatgc	ctagtttcat	ggatgccagc	tggagagcac
14041	cactecenet	agacacttac	agaggagg	yyarcccaag	agcagccatg	gggtgagccc
14101	CCagccaaat	gacaccagag	acayyggaga	catgtgctgc	ggtctgggaa	atagctaccc
14161	aatcgggtcg	catgaaagag	ccattaaaca	ccgcactata	caacatactt	aacttaaacc
14221	CCCCatccc	ctcagcaaaa	yagagagaac	accagtccaa	acagtgcagc	agacccagtt
14281	CCacccccc	gagaagtgcg	cagcagtgtg	gggagctgga	gctggggtgg	ctgtcctgca
14341	ctctanata-	Cgaccctcag aacggctgcc	accacaggca	ctgccaagag	ggaacatgaa	cctagccggc
14401	CCacctatgege	aacggctgcc	cctgacaggt	ggtgacagat	attttcaaga	gtgactctga
14461	totoot	tttccacctt	acatgttgtc	tttggatcct	ttccctgaat	gatatgagat
14521	-grycrggga	actctagccc	tctgtgtgct	gacctccaga	atctgacaac	tttcctttcc
4-J61	uaacagttca	agcaccaagg	cacgatcaca	gtgaacgagg	aaggcaccca	agccaccact
						5

14581	atanaa					
	3-3	tggggttcat	gccgctgtcc	acccaagtcc	gcttcactgt	cgaccgcccc
14641		tcatctacga	gcatcgcacc	agctgcctgc	tcttcatggg	aagagtggcc
14701	aaccccayca	ggtcctagag	gtggaggtct	aggtgtctga	agtgccttgg	ggggacctc
14761	attttgtttc	cattccaaca	acqaqaacaq	agatottoto	cetcetete	cataatttac
14821	gctaccaatc	tgaattcgag	gcccatatga	gaggagctta	geaccacca	cgcagcccac
14881	cttottogaa	tcaattctgc	acaataggg	ataatataa	gaaacyacca	ayaayayayy
14941	cttgttggaa	ctactatta	ataatageee	acyccgcaag	ctcatagaag	tcactgtaac
15001	tgtagtgtgt	ctgctgttat	ctagagggcc	tcacctcccc	actcttcaca	gcaaacctga
10001	gcagcgcgcc	CLAAGCACCE	cccactccaa	tgaccccatc	cttocacacc	taactctatc
12001	acttaagett	LLCLCCacca	ggcccctcat	ctgaatacca	agcacagaaa	trantratat
13121	gactaattee	ttacctctcc	caaggagggt	acacaactag	caccattctt	gatgtccagg
13101	gaagaagcca	ccicaagaca	tatgaggggt	accetagact	aatottaooo	cttaattttc
15241	tcaaagcctg	acctttcaaa	tccatgatga	atoccatcao	teesteetee	tattacatac
15301	ctgtgacctg	gaggacagtg	tataccatat	ctcccatcag	Sessions	cyclycccc
15361	acatttacto	totatotot	ataattatat	cttttatatt	agagataaat	aaatgtagee
15/21	acatttactg	stantone	acaactetet	attttttgaa	gctcaaatat	caaaagccaa
15401	atccaaattc	Ciggataact	ccaggtatga	taaaggctga	gaggaagtca	cttgagcacc
17401	acaacycyce	acagcagggc.	atgitcicaq	gacaggacag	atatatacta	aateetgggg
TODAT	agggreegeg	cagtacccca	gaactgtggg	gtgctaagtg	gcacacaage	cccadaactc
13001	ccacagicia	igccaggctg	ctgcagcttt	cateceteat	acctddtcct	acaataaate
15661	tggtttgaca	gagcagatga	cacctgagga	atatotttct	ggatggttca	atocotocot
15721	aagacaagtg	aaatccacag	aggetattea	acacacaca	ggacccccca	acccctgggc
15781	aggggatgac	tgacggtcac	anntactata	tatassassa	gryceagege	cettecageg
15841	aggggatgac ctggcagat	-gaoggeeae	aggegetgtg	rgrgcaggtg	tctaactgta	accccacage
T2041	ceggeagat					

```
HUMTHRR
                            3472 bp
                                        mRNA
                                                           PRI
                                                                      10-OCT-1991
DEFINITION Human thrombin receptor mRNA, complete cds.
ACCESSION
             M62424
NID
             q339676
KEYWORDS
             thrombin receptor.
SOURCE
             Human DNA.
  ORGANISM Homo sapiens
             Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
             Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
             1 (bases 1 to 3472)
  AUTHORS
             Vu, T.H., Hung, D.T., Wheaton, V.I. and Coughlin, S.R.
             Molecular cloning of a functional thrombin receptor reveals a
  TITLE
novel
             proteolytic mechanism of receptor activation
  JOURNAL
             Cell 64, 1057-1068 (1991)
  MEDLINE
             91168254
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LFVPSVYTGVFVVSLPLNIMAIVVFILKMKVKKPAVVYMLHLATADVLFVSVLPFKIS
YYFSGSDWQFGSELCRFVTAAFYCNMYASILLMTVISIDRFLAVVYPMQSLSWRTLGR
ASFTCLAIWALAIAGVVPLVLKEQTIQVPGLNITTCHDVLNETLLEGYYAYYFSAFSA
VFFFVPLIISTVCYVSIIRCLSSSAVANRSKKSRALFLSAAVFCIFIICFGPTNVLLI
AHYSFLSHTSTTEAAYFAYLLCVCVSSISSCIDPLIYYYASSECQRYVYSILCCKESS
                       DPSSYNSSGQLMASKMDTCSSNLNNSIYKKLLT
BASE COUNT
                           817 c
                                     785 q
                                               937 t
ORIGIN
        61 getegeegag ggtegettgg accetgatet taccegtggg caccetgege tetgeetgee
      121 gegaagaceg geteecegae eegcagaagt eaggagagag ggtgaagegg ageageeega
181 ggeggggeag eeteeeggag eagegeegeg eagageeegg gacaatgggg eegeggegge
      241 tgctgctggt ggccgcctgc ttcagtctgt gcggcccgct gttgtctgcc cgcacccggg
      301 cccgcaggcc agaatcaaaa gcaacaaatg ccaccttaga tccccggtca tttcttctca
      361 ggaaccccaa tgataaatat gaaccatttt gggaggatga ggagaaaaat gaaagtgggt
      421 taactgaata cagattagte tecateaata aaagcagtee tetteaaaaa caactteetg
481 catteatete agaagatgee teeggatatt tgaccagete etggetgaca etetttgtee
      541 catctgtgta caccggagtg tttgtagtca gcctcccact aaacatcatg gccatcgttg
      601 tgttcatcct gaaaatgaag gtcaagaagc cggcggtggt gtacatgctg cacctggcca
      661 cggcagatgt gctgtttgtg tctgtgctcc cctttaagat cagctattac ttttccggca
      721 gtgattggca gtttgggtct gaattgtgtc gcttcgtcac tgcagcattt tactgtaaca
781 tgtacgcctc tacttgctc atgacagtca taagcattga ccggtttctg gctgtggtgt
841 atcccatgca gtccctctcc tggcgtactc tgggaagggc ttccttcact tgtctggcca
      901 tetgggettt ggecategea ggggtagtge etetegteet caaggageaa accatecagg
      961 tgcccgggct caacatcact acctgtcatg atgtgctcaa tgaaaccctg ctcgaaggct
     1021 actatgccta ctacttctca gccttctctg ctgtcttctt ttttgtgccg ctgatcattt
     1081 ccacggtctg ttatgtgtct atcattcgat gtcttagctc ttccgcagtt gccaaccgca
     1141 gcaagaagte cegggetttg tteetgteag etgetgttt etgeatette atcatttget
1201 teggacecae aaacgteete etgattgege attacteatt cettteteac aetteeacea
    1261 cagaggetge ctactttgcc tacctcctct gtgtctgtgt cagcagcata agctcgtgca
    1321 tegacecect aatttactat tacgetteet etgagtgeea gaggtacgte tacagtatet
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						•	
13	381	tatgctgcaa	agaaagttcc	gatcccagca	gttataacag	cagtggggag	ttgatggcaa
-		geddddelggd	Caccicici	agraaccroa	araananat	2 t 2 0 2 2 2 2 2 2	atattaaatt
		aggaaaagggg	actyctygga	ggttaaaaag	aaaagtttat	aaaagtgaat	aacctgagga
		Lectateage	ccccacccaa	actttattga	ttcacctcct	aaaacaacad	atotacoact
		egeatacety	ctttttatgg	gagetgteaa	gcatgtattt	ttatcaatta	ccagaagat
	581	uacaggacga	gatgacggtg	ttattccaag	ggaatattgc	caatoctaca	ataataaata
	741	auch carre	CLYGALALAG	CLAGGEGACA	ratacatact	tacate**	tatatataaa
18	301	eguatguata	Cacatatatt	atttgcagtg	Cartataraa	t=~~~~~+++	2222222
		cccccgcac	CCCaycaatt	atdaaaaraa	Tetetastte	CCtcatttaa	Fatocasact
		ccaggiciggi	agagillage	cctdaacatt	tcataatatt	catcaacact	Managage ag
		acagectggg	CLLYLACIAC	LLLEGCAAAF	aaatatatt	+ ~ = = = + + + + + + + + + + + + + + +	tanaaaaaa
	, 11	geecaageea	LLaayayyta	adacttagra	ctatctotoc	atamaaatta	tantattta
		uacccaaac	atatecaage	LLUAALLCCL	aaaattataa	aaacadatka	2220001010
		cccegacacg	ggtagtatt	LLLacattrr	acacactota.	Cacataacco	22224ta2aa
22	221	ataagtcctc	tagtgaatgt	aggctggctt	tcacactaca	Ctattcotca	aaaactyayc
22	281	tgtccgcccc	cqatqqaqqa	ctccaggcag	cadageagg	ccacccccga	gagetgeatg
23	341	gattggccag	aaaccttcct	gctgagcctc	acaccacacg	ccayyyccat	gudagadada
24	101	ctccatcctc	ctgggattgg	ctgtgaactg	atcatottta	taacaaaata	actacattty
24	61	atgtgatatc	ctaggaggta	atgaccatga	accacyccta	taggaaactg	gcaaagcaga
~ -		aaagaaggca	Lydacticia	gatgcccarc	Cactgggtgt	2220202101	~~+~~++~+ <i></i>
25	81	ctgaaatgtc	agttctgata	tggaagcacc	cattatococ	tatagaaaat	agragitytt
		cegagegeae	agautuuaat	aadacadada	CCtaccctca	2727722277	20210210
27	701	tagagtgtga	tgtatgtgta	ataaatatgt	treacacasa	caaccacat	agattatgea
27	61	agtttgaaca	tttgggttac	tatttcttgt	oottataact	taatgaaaa	cagetaaaga
28	321	aggacatata	ttttttaaaa	taagtctgat	ttaattaacc	aatsttatt	aatgeagtae
28	881	ttgctcaata	gattgctcaa	atcaggtttt	cttttaagaa	tanatantat	cacaaatgtt
29	41	agaaataaca	gaagaaaata	gaattgacat	trasatrtar	gazazetet	cagtetgett
30	01	catttactta	agacttaatg	agactttaaa	accettttt	gaaaattatt	ctataatttc
30	61	tagaaaatct	tcatggaart	cacaaagtaa	tttaaaaaat	aacctcctaa	gtatcaagta
31	.21	tcttacgaaa	aaatggtagc	attttaaaca	anatagaaatt	aggttgaaac	ataccccta
31	81	taaaagagca	daccadacac	ggtggctcac	adatayaday	Lugcaaggea	aatgtttatt
32	41	ggcgggtgga	tcacgaggtc	aggagatcga	geergeate	ccagcacttt	gggaggctga
33	01	ctctactaaa	aatgcaaaaa	agattacco	gaccateetg	getaacacgg	tgaaacccgt
33	61	tactcgggag	actaaaacaa	aaattagccg gagactggcg	ggcgcgggcgg	caggcacctg	tagtcccagc
34	21	cgagatcgcg	ccactatact	ccagcctggg	cyaacccagg	aggeggaeet	tgtagtgagc
		5509		ccagcccggg	caacagagca	agactccatc	CC

```
LOCUS
            HUMLPLFI
                          3877 bp
                                      DNA
                                                       PRI
                                                                 07-JAN-1995
DEFINITION
            H. sapiens lipoprotein lipase (LPL) gene, exons 7,8, and 9, and an
            Alu repetative element.
ACCESSION
            M76722 M76723
NID
            g187215
KEYWORDS
            Alu repeat; lipoprotein lipase; plasma protein.
SOURCE
            Homo sapiens blood DNA.
  ORGANISM Homo sapiens
            Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata;
            Vertebrata; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 3877)
REFERENCE
  AUTHORS
            Chuat, J.C., Raisonnier, A., Etienne, J. and Galibert, F.
  TITLE
            The lipoprotein lipase-encoding human gene: sequence from intron-6
            to intron-9 and presence in intron-7 of a 40-million-year-old Alu
            sequence
  JOURNAL
            Gene 110 (2), 257-261 (1992)
  MEDLINE
            92165069
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                      /tissue_type="blood"
                      /map="8p22"
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                      /partial
                      /gene="LPL"
                      /note="G00-120-700"
                      /number=6
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translation="FHYQVKIHFSGTESETHTNQAFEISLYGTVAESENIPFTLPEVS/
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                      IFCSREKVSHLQKGKAPAVFVKCHDKSLNKKSG*
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                      /number=7
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FIG. 36A

ez:-" . .

93/97

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                      /partial
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                     /note="G00-120-700"
                     /number=9
BASE COUNT
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                                   746 g
                                           1199 t
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       61 acaccagtgg ttccatgtgt gtgcacttcc ggtttgagtg ctagtgagat acttctgtgg
      121 ttctgaattg cctgactatt tggggttgtg atattttcat aaagattgat caacatgttc
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      241 gtgaaaccca taccaatcag gcctttgaga tttctctgta tggcaccgtg gccgagagtg
      301 agaacateee atteactetg tgagtageac aggggggegg teateatgge accagteeet
     361 ctcctgccat aacccttggt ctgagcagca gaagcagaga gcgatgccta gaaaacaagt
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     541 tettatatet gettatetet tetataaage tgetgetaaa caatataate aaactatete
     601 aaaaggtttg acattaaaga aaatgagcaa tggtaacagg aaaccactct atagatgtac
     661 atataatatg tacagaaaat ataagtagta agaagtccat gacaaagtgt tagctctttt
     721 ttttttttt tttttttt tttttgagat ggagtetete tetattgeec aggetggagt
     781 gcagtgattc gatctcagct cactgcaacc tctacctccc gagttcaaac aattcttctg
     841 tetcageete eegagtaget ggggetgeag gtgcccacca ccatgeecag etaatttttg
     901 tattttagt agcgacaggg teteaceatg ttggccaage tggtettgaa tteetgatet
     961 caggtgatec accegeeteg geeteccaaa gtgetgggat tacaggtgtg agceaccatg
    1021 cccagcctac cctttactac taatcaaaga aataaaagta aggcaacttg atacttttac
    1081 aattactaga tgaacaaatc tttaaaaata gccagtgcag acaaggtggt gaagcagaac
    1141 atgcgaacct accatgcatc attcacggct agaaccetce aggtgcggaa ggtagtattt
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                 (bases 1 to 2566)
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              Calvo, D. and Vega, M.A.
              Identification, primary structure, and distribution of CLA-1, a novel member of the CD36/LIMPII gene family
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              J. Biol. Chem. 268 (25), 18929-18935 (1993)
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# **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



# INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

51) International Patent Classification ⁶ :		(11) International Publication Number: WO 99/50454
C12Q 1/68	A3	(43) International Publication Date: 7 October 1999 (07.10.99
21) International Application Number: PCT/US  22) International Filing Date: 26 March 1999 (		& Reynolds, P.C., Two Militia Drive, Lexington, MA 0242
(30) Priority Data: 09/054,272 1 April 1998 (01.04.98)  (63) Related by Continuation (CON) or Continuation-in (CIP) to Earlier Application US 09/054, Filed on 1 April 1998 (  (71) Applicant (for all designated States except US): WHI INSTITUTE FOR BIOMEDICAL RESEARCH Nine Cambridge Center, Cambridge, MA 02142 (  (72) Inventors; and ((75) Inventors/Applicants (for US only): LANDER, [US/US]; 151 Bishop Allen Drive, Cambridge, M (US). DALEY, George, Q. [US/US]; 50 You Weston, MA 02193 (US). CARGILL, Michele 50 Follen Street #208, Cambridge, MA 021 IRELAND, James, S. [US/US]; 36 College A' Somerville, MA 02144 (US). ROZEN, Steven, G. 45 Josephine Avenue, Somerville, MA 02144-23	UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MI RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DE ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MI NE, SN, TD, TG).  Published  With international search report.  Before the expiration of the time limit for amending the claim and to be republished in the event of the receipt of amendments and to be republication of the international search report:  [88] Date of publication of the international search report:  13 April 2000 (13.04.0	
of a gene, including polymorphic sites. Allele-specific 1	the hun	CULAR PATHOLOGY GENES  an genome, particularly nucleic acid segments from the coding regi and probes hybridizing to regions flanking or containing these sites a applications such as phenotype correlations, forensics, paternity testing

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Interna .al Application No PCT/US 99/06473

		1 101	/05 99/064/3						
A. CLASSI IPC 6	A. CLASSIFICATION OF SUBJECT MATTER  I PC 6 C12Q1/68								
According to	o International Patent Classification (IPC) or to both national classifica	tion and IPC							
	SEARCHED								
IPC 6	ocumentation searched (classification system followed by classification C12Q	n symbols)							
Documental	tion searched other than minimum documentation to the extent that $oldsymbol{s}$ .	uch documents are included in th	e fields searched						
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)									
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT								
Category *	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.						
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X	1-4,11,								
Y									
1	see abstract		10						
X	1-4,11,								
Υ	XP002121302 see abstract 		10						
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اختا	her documents are listed in the continuation of box C.	X Patent family members	are listed in annex.						
"A" docume consider filing de "L" docume which citation "O" docume other i "P" docume "P" docume "P" docume i	tegories of cited documents:  ent defining the general state of the art which is not letted to be of particular relevance to current but published on or after the international late with which may throw doubts on priority claim(s) or is called to establish the publication date of another nor other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means art published prior to the international filing date but has the priority date claimed.	er the international filing date unflict with the application but ciple or theory underlying the ance; the claimed invention or cannot be considered to her the document is taken alone ance; the claimed invention roive an inventive step when the one or more other such docu- eing obvious to a person skilled							
<del></del>	actual completion of the international search	*&* document member of the sa Date of mailing of the intern							
	November 1999		2. 2000						
Name and r	mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Authorized officer							
	Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Knehr, M							

1

Intern. Ial Application No PCT/US 99/06473

ategory °	DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
·		There was to claim No.	
<b>K</b>	DÜRR C ET AL.: "Genetic studies of antithrombin III with IEFand ASO hybridization" HUMAN GENETICS, vol. 90, 1992,	5-7,11, 12	
•	pages 457-459, XP002121293 see the whole document	10	
X	OKAJIMA K ET AL.: "Antithrombin III Nagasaki (Ser116-Pro): A heterozygous variant with defective heparin binding associated with thrombosis" BLOOD, vol. 81, no. 5, 1993,	5-7,11, 12	
Y	pages 1300-1305, XP002121294 see abstract	10	
X	UEYAMA H ET AL.: "Antithrombin III Kumamoto: Identification of a point mutation and genotype analysis of the family" THROMBOSIS AND HAEMOSTASIS, vol. 63, no. 2, 1990,	5-7	
Y	pages 231-234, XP002121295 see the whole document	10-12	
<b>K</b>	ZEE R Y L ET AL.: "Association and linkage analysis of restriction fragment length polymorphisms for the human renin and antithrombin III genes in essential hypertension" JOURNAL OF HYPERTENSION, vol. 9, 1991, pages 825-830, XP002121296 see the whole document	11,12	
(	BOCK S C ET AL.: "Antithrombin III Utah: Proline-407 to leucine mutation in a highly conserved region near the inhibitor reactive site" BIOCHEMISTRY, vol. 27, 1988, pages 6171-6178, XP002121297 cited in the application see the whole document	11,12	
•	BELGRADER P ET AL.: "A multiplex PCR-ligase detection reaction assay for human identity testing" GENOME SCIENCE & TECHNOLOGY, vol. 1, no. 2, 1996, pages 77-87, XP002121298  * see especially Fig. 1 and Table 1 * see the whole document	5,8-12	

Interna. al Application No PCT/US 99/06473

C./Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/US 99/06473	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
Y	SYVANEN A -CH ET AL: "IDENTIFICATION OF INDIVIDUALS BY ANALYSIS OF BIALLELIC DNA MARKERS, USING PCR AND SOLID-PHASE MINISEQUENCING" AMERICAN JOURNAL OF HUMAN GENETICS, vol. 52, no. 1, 1 January 1993, pages 46-59, XP002050638 see the whole document	5,8,9	
1	WO 95 12607 A (MOLECULAR TOOL INC) 11 May 1995 * see especially the claims * see the whole document		
A	WANG D ET AL: "TOWARD A THIRD GENERATION GENETIC MAP OF THE HUMAN GENOME BASED ON BI-ALLELIC POLYMORPHISMS" AMERICAN JOURNAL OF HUMAN GENETICS, vol. 59, no. 4, 1 October 1996, page A03 XP002050641 see abstract		
P,X	WO 98 20165 A (WHITEHEAD BIOMEDICAL INST; HUDSON THOMAS (US); LANDER ERIC S (US);) 14 May 1998 see the whole document	1-12	
P,X	DALEY G Q ET AL.: "High throughput polymorphism discovery in genes related to thrombosis: A paradigm for linking common variants to common disease" BLOOD, vol. 92, no. 10/1, 1998, page 1953 XP002121299 see abstract	11,12	
T	CARGILL M ET AL.: "Characterization of single-nucleotide polymorphisms in coding regions of human genes" NATURE GENETICS, vol. 22, 1999, pages 231-238, XP002121300 see the whole document	1-4, 10-12	

International application No. PCT/US 99/06473

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
Claims Nos.:     because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
see additional sheet, subject 1.
· ·
Remark on Protest  The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

### 1. Claims: 1-12 (partially)

INVENTION 1: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the AT3 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

### 2. Claims: 1-12 (partially)

INVENTION 2: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the CETP gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 3. Claims: 1-12 (partially)

INVENTION 3: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the CLanalog gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 4. Claims: 1-12 (partially)

INVENTION 4: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the F2R gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 5. Claims: 1-12 (partially)

INVENTION 5: A nucleic acid molecule of at least 5

nucleotides in length consisting of a part of the F2 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 6. Claims: 1-12 (partially)

INVENTION 6: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the F3 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

### 7. Claims: 1-12 (partially)

INVENTION 7: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the F5 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

### 8. Claims: 1-12 (partially)

INVENTION 8: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the HCF2 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

# 9. Claims: 1-12 (partially)

INVENTION 9: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the HMGCR gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table

- column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 10. Claims: 1-12 (partially)

INVENTION 10: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the ITGA2B gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

## 11. Claims: 1-12 (partially)

INVENTION 11: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the ITB3 gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

### 12. Claims: 1-12 (partially)

INVENTION 12: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the LCAT gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

# 13. Claims: 1-12 (partially)

INVENTION 13: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the LDLR gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing

such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 14. Claims: 1-12 (partially)

INVENTION 14: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the LPL gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

### 15. Claims: 1-12 (partially)

INVENTION 15: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the PROC gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 16. Claims: 1-12 (partially)

INVENTION 16: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the PTAFR gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

#### 17. Claims: 1-12 (partially)

INVENTION 17: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the TFPI gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

18. Claims: 1-12 (partially)

INVENTION 18: A nucleic acid molecule of at least 5 nucleotides in length consisting of a part of the TBXA2R gene and comprising a polymorphic (biallelic) site according to the nucleotide positions as indicated in the attached table - column 4, an allele-specific oligonucleotide hybridizing to such a polymorphic site, an isolated gene product encoded by such a nucleic acid molecule, and a method of analyzing such a nucleic acid by determining the bases occupying the polymorphic site(s).

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Information on patent family members

Interna. al Application No PCT/US 99/06473

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